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#### TITLE OF THE INVENTION

POLYNUCLEIC ACIDS ISOLATED FROM A PORCINE REPRODUCTIVE AND RESPIRATORY SYNDROME VIRUS (PRRSV), PROTEINS ENCODED BY THE POLYNUCLEIC ACIDS, VACCINES BASED ON THE PROTEINS AND/OR POLYNUCLEIC ACIDS, A METHOD OF PROTECTING A PIG FROM A PRRSV AND A METHOD OF DETECTING A PRRSV

This is a continuation-in-part of application Serial No. 08/131,625, filed on October 5, 1993, pending, which is a continuation-in-part of application Serial No. 07/969,071, filed on October 30, 1992, now abandoned. The entire contents of application Serial No. 08/131,625, filed on October 5, 1993, are incorporated herein by reference.

### BACKGROUND OF THE INVENTION

#### Field of the Invention

The present invention concerns DNA isolated from a porcine reproductive and respiratory virus (PRRSV), a protein and/or a polypeptide encoded by the DNA, a vaccine which protects pigs from a PRRSV based on the protein or DNA, a method of protecting a pig from a PRRSV using the vaccine, a method of producing the vaccine, a method of treating a pig infected by or exposed to a PRRSV, and a method of detecting a PRRSV.



In recent years, North American and European swine herds have been susceptible to infection by new strains of reproductive and respiratory viruses (see A.A.S.P., September/October 1991, pp. 7-11; The Veterinary Record, February 1, 1992, pp. 87-89; Ibid., November 30, 1991, pp. 495-496; Ibid., October 26, 1991, p. 370; Ibid., October 19, 1991, pp. 367-368; Ibid., August 3, 1991, pp. 102-103; Ibid., July 6, 1991; Ibid., June 22, 1991, p. 578; Ibid., June 15, 1991, p. 574; Ibid., June 8, 1991, p. 536; Ibid., June 1, 1991, p. 511; Ibid., March 2, 1991, p. 536; Ibid., June 1, 1991, p. 511; Ibid., March 2, 1991, p. 213). Among the first of the new strains to be identified was a virus associated with the so-called Mystery Swine Disease (MSD) or "blue-eared syndrome", now known as Swine Infertility and Respiratory Syndrome (SIRS) or Porcine Reproductive and Respiratory Syndrome (PRRS).

An MSD consisting of reproductive failure in females and respiratory disease in nursing and weaned pigs appeared in the midwestern United States in 1987 (Hill et al., Am. Assoc. Swine Practitioner Newsletter 4:47 (1992); Hill et al., Proceedings Mystery Swine Disease Committee Meeting, Denver, Colorado 29-31 (1990); Keffaber, Am. Assoc. Swine Practitioner Newsletter 1:1-9 (1989); Loula, Agri-Practice 12:23-34 (1991)). Reproductive failure was characterized by abortions, stillborn and weak-born pigs. The respiratory disease in nursing and weaned pigs was

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characterized by fever, labored breathing and pneumonia. A similar disease appeared in Europe in 1990 (Paton et al., Vet. Rec. 128:617 (1991); Wensvoort et al., Veterinary Quarterly 13:121-130 (1991); Blaha, Proc. Am. Assoc. Swine Practitioners, pp. 313-315 (1993)), and has now been recognized worldwide.

This disease has also been called porcine epidemic abortion and respiratory syndrome (PEARS), blue abortion disease, blue ear disease (U.K.), abortus blau (Netherlands), seuchenhafter spatabort der schweine (Germany), Heko-Heko disease, and in the U.S., Wabash syndrome, mystery pig disease (MPD), and swine plague (see the references cited above and Meredith, Review of Porcine Reproductive and Respiratory Disease Syndrome, Pig Disease Information Centre, Department of Veterinary Medicine, Madingley Road, Cambridge CB3 OES, U.K. (1992); Wensvoort et al., Vet. Res. 24:117-124 (1993); Paul et al., J. Clin. Vet. Med. 11:19-28 (1993)). In Europe, the corresponding virus has been termed "Lelystad virus."

At an international conference in May, 1992, researchers from around the world agreed to call this disease Porcine Reproductive and Respiratory Syndrome (PRRS). The disease originally appeared to be mainly a reproductive disease during its early phases, but has now evolved primarily into a respiratory disease.

Porcine reproductive and respiratory syndrome virus (PRRSV) is a relatively recently recognized swine pathogen associated with porcine reproductive and respiratory syndrome (PRRS). PRRSV is a significant pathogen in the swine industry. PRRSV infections are common in the U.S. swine herds. Outbreaks of PRRS in England have led to cancellation of pig shows.

The symptoms of PRRS include a reluctance to eat (anorexia), a mild fever (pyrexia), cyanosis of the extremities (notably bluish ears), stillbirths, abortion, high mortality in affected litters, weak-born piglets and premature farrowing. The majority of piglets born alive to affected sows die within 48 hours. PRRS clinical signs include mild influenza-like signs, rapid respiration ("thumping"), and a diffuse interstitial pneumonitis. virus has an incubation period of about 1-2 weeks from The virus appears to contact with a PRRSV-infected animal. be an enveloped RNA arterivirus (The Veterinary Record, February 1, 1992). The virus has been grown successfully in pig alveolar macrophages and CL2621 cells (Benfield et al, J. Vet. Diagn. Invest., 4:127-133, 1992; Collins et al, Swine Infertility and Respiratory Syndrome/Mystery Swine Disease. Proc., Minnesota Swine Conference for Veterinarians, pp. 200-205, 1991), and in MARC-145 cells (Joo, PRRS: Diagnosis, Proc., Allen D. Leman Swine Conference, Veterinary Continuing Education and Extension,

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University of Minnesota (1993), 20:53-55; <u>Kim et al</u>, Arch. Virol., 133:477-483 (1993)). A successful culturing of a virus which causes SIRS has also been reported by <u>Wensvoort et al</u> (Mystery Swine Disease in the Netherlands: The Isolation of Lelystad Virus. Vet. Quart. 13:121-130, 1991).

Initially, a number of agents were incriminated in the etiology of this disease (Wensvoort et al., Vet. Res. 24:117-124 (1993); Woolen et al., J. Am. Vet. Med. Assoc. 197:600-601 (1990)). There is now a consensus that the causative agent of PRRS is an enveloped RNA virus referred to as Porcine Reproductive and Respiratory Syndrome Virus (PRRSV), reportedly of approximately 62 nm in diameter (Benfield et al., J. Vet. Diagn. Invest., 4:127-133, 1992).

Virus isolates vary in their ability to replicate in continuous cell lines. Some grow readily, while others require several passages and some grow only in swine alveolar (SAM) cultures (Bautista et al., J. Vet. Diagn. Invest. 5:163-165, 1993; see also the Examples hereunder [particularly Table 1]).

PRRSV is a member of an Arterivirus group which includes equine arteritis virus (EAV), lactate dehydrogenase-elevating virus (LDV) and simian hemorrhagic fever virus (SHFV) (Benfield et al., 1992, supra; Plagemann, Proc. Am. Assoc. Swine Practitioners, 4:8-15 1992; Plagemann and Moennig, Adv. Virus Res. 41:99-192,

1992; Conzelmann et al., Virology, 193:329-339, 1993;

Godney et al., Virology, 194:585-596, 1993; Meulenberg et al., Virology, 192:62-72, 1993). The positive-strand RNA viruses of this Arterivirus group resemble togaviruses morphologically, but are distantly related to coronaviruses 5 and toroviruses on the basis of genome organization and gene expression (Plagemann et al., supra; Spaan et al., J. Gen. Virol. 69, 2939-2952 (1988); Strauss et al., Annu. Rev. Biochem. 42, 657-683 (1988); Lai, Annu. Rev. 10 Microbiol. 44, 303-333 (1990); Snijder et al., Nucleic Acid Res. 18, 4535-4542 (1990)). The members of this group infect macrophages and contain a nested set of 5 to 7 subgenomic mRNAs in infected cells (Plagemann et al., supra; Meulenberg et al., Virology, 192, 62-72 (1993); ត្រូ 15 Conzelmann et al., Virology, 193, 329-339 (1993); 15, 16, 17, 18, 19).

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The viral genome of European isolates has been shown to be a plus stranded RNA of about 15.1 kb (Conzelmann et al., supra; Meulenberg et al., supra), and appears to be similar in genomic organization to LDV and EAV (Meulenberg et al., supra). However, no serological cross-reaction has been found among PRRSV, LDV and EAV (Goyal et al., J. Vet. Diagn. Invest., 5, 656-664 (1993)).

PRRSV was initially cultivated in swine alveolar macrophage (SAM) cell cultures (Pol et al., Veterinary Quarterly, 13:137-143, 1991; Wensvoort et al., Veterinary

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Quarterly, 13:121-130, 1991) and then in continuous cell lines CL2621 (Benfield et al., supra), MA-104, and MARC-145 (Joo, Proc. Allen D. Leman Swine Conference, pp. 53-55, 1993). The reproductive and respiratory disease has been reproduced with cell free lung filtrates (Christianson et al., Am. J. Vet. Res., 53:485-488, 1992; Collins et al., J. Vet. Diagn. Invest., 4:117-126, 1992; Halbur et al., Proc. Central Veterinary Conference, pp. 50-59, 1993), and with cell culture-propagated PRRSV (Collins et al., supra, and Proc. Allen D. Leman Swine Conference, pp. 47-48, 1993).

Eight open reading frames (also referred to herein as "ORFs" or "genes") have been identified in a European PRRSV isolate. The genes of this European isolate are organized similarly to that in coronavirus (Meulenberg et al., supra). A 3'-end nested set of messenger RNA has been found in PRRSV-infected cells similar to that in coronaviruses (Conzelmann et al., supra; Meulenberg et al., supra).

The ORF 1a and 1b at the 5'-half of the European PRRSV genome are predicted to encode viral RNA polymerase. The ORF's 2-6 at the 3'-half of the genome likely encode for viral membrane-associated (envelope) proteins (Meulenberg et al., supra). ORF6 is predicted to encode the membrane protein (M) based on its similar characteristics with the ORF 6 of EAV, ORF 2 of LDV, and the M protein of mouse hepatitis virus and infectious bronchitis virus (Meulenberg

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et al., Virology 192, 62-72 (1993); Conzelmann et al.,
Virology 193, 329-339 (1993); Murtaugh, Proc. Allen D.

Leman Swine Conference, Minneapolis, MN, pp. 43-45 (1993);

Mardassi et al., Abstracts of Conference of Research

Workers in Animal Diseases, Chicago, IL, pp. 43 (1993)).

The product of ORF 7 is extremely basic and hydrophilic,
and is predicted to be the viral nucleocapsid protein (N)

(Meulenberg et al., supra; Conzelmann et al., supra;

Murtaugh, supra; Mardassi et al., supra and J. Gen. Virol.,
75:681-685 (1994)).

Although conserved epitopes have been identified between U.S. and European PRRSV isolates using monoclonal antibodies (Nelson et al., J. Clin. Microbiol., 31:3184-3189, 1993), there is extensive antigenic and genetic variation both among U.S. and European isolates of PRRSV (Wensvoort et al., J. Vet. Diagn. Invest., 4:134-138, 1992). European isolates are genetically closely related, as the nucleotide sequence at the 3'-half of the genome from two European PRRSV isolates is almost identical (Conzelmann et al., supra; Meulenberg et al., supra).

Although the syndrome caused by PRRSV appears to be similar in the U.S. and Europe, several recent studies have described phenotypic, antigenic, genetic and pathogenic variations among PRRSV isolates in the U.S. and in Europe (Murtaugh, supra; Bautista et al., J. Vet. Diagn. Invest., 5, 163-165 (1993); Bautista et al., J. Vet. Diagn. Invest.,

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5, 612-614 (1993); Wensvoort/et al., J. Vet. Diagn. Invest., 4, 134-138 (1992); Stevenson et al., J. Vet. Diagn. Invest., 5, 432-434 (1993)). For example, the European isolates grow preferentially in SAM cultures and replicate to a very low titer in other culture systems (Wensvoort, Vet. Res., 24, 117-124 (1993); Wensvoort et al., J. Vet. Quart., 13, 121-130 (1991); Wensvoort et al., J. Vet. Diagn. Invest., 4, 134-138 (1992)). On the other hand, some of the U.S. isolates have been shown to replicate well in SAM as well as in the continuous cell line CL2621 (Benfield et al., J. Vet. Diagn. Invest., 4, 127-133 (1992); Collins et al., J. Vet. Diagn. Invest., 4, 117-126 (1992)). Thus, phenotypic differences among U.S. isolates are observed, as not all PRRSV isolates isolated on SAM can replicate on the CL2621 cell line (Bautista et al., J. Vet. Diagn. Invest., 5, 163-165 (1993)).

A high degree of regional antigenic variation among PRRSV isolates may exist. Four European isolates were found to be closely related antigenically, but these European isolates differed antigenically from U.S. isolates. Further, three U.S. isolates were shown to differ antigenically from each other (Wensvoort et al., J. Vet. Diagn. Invest., 4, 134-138 (1992)). Animals seropositive for European isolates were found to be negative for U.S. isolate VR 2332 (Bautista et al., J. Vet. Diagn. Invest., 5, 612-614 (1993)).

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U.S. PRRSV isolates differ genetically at least in part from European isolates (Conzelmann et al., supra; Meulenberg et al., supra; Murtaugh et al., Proc. Allen D. Leman Conference, pp. 43-45, 1993). The genetic differences between U.S. and European isolates are striking, especially since they are considered to be the same virus (Murtaugh, supra). Similar observations were also reported when comparing the Canadian isolate IAF-exp91 and another U.S. isolate VR 2332 with LV (Murtaugh, supra; Mardassi, supra). However, the 3' terminal 5 kb nucleotide sequences of two European isolates are almost identical (Conzelmann et al., supra; Meulenberg et al., supra).

The existence of apathogenic or low-pathogenic strains among isolates has also been suggested (Stevenson, supra). Thus, these studies suggest that the PRRSV isolates in North America and in Europe are antigenically and genetically heterogeneous, and that different genotypes or serotypes of PRRSV exist. However, prior to the present invention, the role of antigenic and genetic variation in the pathogenesis of PRRSV was not entirely clear.

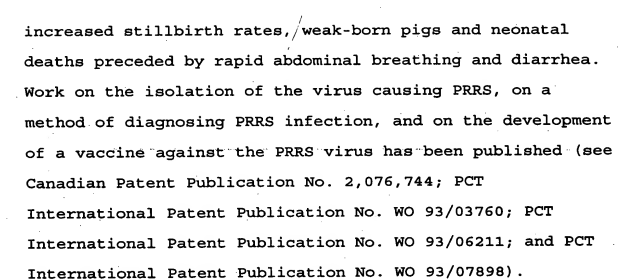
The occurrence of PRRS in the U.S. has adversely affected the pig farming industry. Almost half of swine herds in swine-producing states in the U.S. are seropositive for PRRSV (Animal Pharm., 264:11 (11/11/92)). In Canada, PRRS has been characterized by anorexia and pyrexia in sows lasting up to 2 weeks, late-term abortions,

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There is also variability in the virulence of PRRSV in herds. Recently, a more virulent form of PRRS has been occurring with increased incidence in 3-8 week old pigs in the midwestern United States. Typically, healthy 3-5 week old pigs are weaned and become sick 5-7 days later.

Routine virus identification methods on tissues from affected pigs have shown that swine influenza virus (SIV), pseudorabies virus (PRV), and Mycoplasma hyopneumoniae are not associated with this new form of PRRS. Originally termed proliferative interstitial pneumonia (PIP; see U.S. patent application Serial No. 07/969,071), this disease has been very recently linked with PRRS, and the virus has been informally named the "Iowa strain" of PRRSV (see U.S. patent application Serial No. 08/131,625).

Pessimism and skepticism has been expressed in the art concerning the development of effective vaccines against these porcine viruses (The Veterinary Record, October 26,

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1991). A belief that human influenza vaccine may afford some protection against the effects of PRRS and PNP exists (The Veterinary Record, July 6, 1991).

Viral envelope proteins are known to be highly variable in many coronaviruses, such as feline infectious peritonitis virus and mouse hepatitis virus (Dalziel et al: Site-specific alteration of murine hepatitis virus type 4 peplomer glycoprotein E2 results in reduced neurovirulence. J. Virol., 59:464-471 (1986); Fleming et al: Pathogenicity of antigenic variants of murine coronavirus JHM selected J. Virol., 58:869-875 (1986); with monoclonal antibodies. Fiscus et al: Antigenic comparison of the feline coronavirus isolates; Evidence for markedly different peplomer glycoproteins. J. Virol., 61:2607-2613 (1987); Parker et al: Sequence analysis reveals extensive polymorphism and evidence of deletions within the E2 glycoprotein gene of several strains of murine hepatitis virus. Virology, 173:664-673 (1989)).

For example, a deletion or a mutation in the major envelope protein in coronaviruses can alter tissue tropism and in vivo pathogenicity. A mutation in a monoclonal antibody-resistant mutant of MHV has resulted in loss of its neurovirulence for mice (Fleming et al, 1986 supra). Porcine respiratory coronavirus (PRCV) is believed to be a deletion mutant of transmissible gastroenteritis virus (TGEV) in swine. The deletion in the PRCV genome may be in

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the 5'-end of the spike (S) gene of TGEV (Halbur et al, An overview of porcine viral respiratory disease. Proc.

Central Veterinary Conference, pp. 50-59 (1993); Laude et al, Porcine respiratory coronavirus: Molecular features and virus-host interactions. Vet. Res., 24:125-150 (1993); Vaughn et al, Isolation and characterization of three porcine respiratory coronavirus isolates with varying sizes of deletions. J. Clin. Micro., 32:1809-1812 (1994)).

PRCV has a selective tropism for the respiratory tract and does not replicate in the gastrointestinal tract (Rasschaert et al, Porcine respiratory coronavirus differs from transmissible gastroenteritis virus by a few genomic deletions. J. Gen. Virol., 71:2599-2607 (1990); Laude et al, 1993 supra). In contrast, TGEV has a tropism for both respiratory and gastrointestinal tracts (Laude et al, 1993 supra).

Variation in antigenic and genetic relatedness among
LDV isolates of varying pathogenicity is also known (Kuo et
al, Lactate-dehydrogenase-elevating virus (LDV):

20 subgenomic mRNAs, mRNA leader and comparison of 3'-terminal
sequences of two LDV isolates. Virus Res., 23:55-72
(1992); Plagemann, LDV, EAV, and SHFV: A new group of
positive stranded RNA viruses. Proc. Am. Assoc. Swine
Practitioners, 4:8-15 (1992); Chen et al, Sequences of 3'
25 end of genome and of 5' end of open reading frame 1a of
lactate dehydrogenase-elevating virus and common junction



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motifs between 5' leader and bodies of seven subgenomic mRNAs. J. Gen. Virol., 74:643-660 (1993)).

However, the present invention provides the first insight into the relationships between the open reading frames of the PRRSV genome and their corresponding effects on virulence and replication.

Further, a diagnosis of porcine reproductive and respiratory syndrome (PRRS) relies on compiling information from the clinical history of the herd, serology, pathology, and ultimately on isolation of the PRRS virus (PRRSV). Three excellent references reviewing diagnosis of PRRSV have been published in the last year (Van Alstine et al, "Diagnosis of porcine reproductive and respiratory syndrome, " Swine Health and Production, Vol. 1, No. 4 (1993), p. 24-28; Christianson et al, "Porcine reproductive and respiratory syndrome: A review." Swine Health and Production, Vol. 1, No. 2 (1994), pp. 10-28 and Goyal, "Porcine reproductive and respiratory syndrome, " J. Vet. Diagn. Invest. 5:656-664 (1993)). PRRSV has also recently been shown to replicate in pulmonary alveolar macrophages by gold colloid immunohistochemistry (Magar et al (1993): Immunohistochemical detection of porcine reproductive and respiratory syndrome virus using colloidal gold. Can. J. Vet. Res., 57:300-304).

Clinical signs vary widely between farms, and thus, are not the most reliable evidence of a definitive

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diagnosis, except in the case of a severe acute outbreak in naive herds which experience abortion storms, increased numbers of stillborn pigs, and severe neonatal and nursery pig pneumonia. Presently, the most common clinical presentation is pneumonia and miscellaneous bacterial problems in 3-10 week old pigs. However, many PRRSV-positive herds have no apparent reproductive or respiratory problems.

Some herds evidence devastating reproductive failure, characterized by third-trimester abortions, stillborn pigs and weak-born pigs. Many of these herds also experience severe neonatal respiratory disease. Respiratory disease induced by PRRSV in 4-10 week-old pigs is also common and can be quite severe (Halbur et al, Viral contributions to the porcine respiratory disease complex. Proc. Am. Assoc. Swine Pract. (1993), pp. 343-350). Clinical PRRSV outbreaks are frequently followed by bacterial pneumonia, septicemia, or enteritis. Thus, it has been difficult to obtain an acceptably rapid and reliable diagnosis of infection by PRRSV, prior to the present invention.

The pig farming industry has been and will continue to be adversely affected by these porcine reproductive and respiratory diseases and new variants thereof, as they appear. PRRSV is a pathogen of swine that causes economic losses from reproductive and respiratory diseases.

Economic losses from PRRS occur from loss of pigs from

abortions, stillborn pigs, repeat breeding, pre-weaning and postweaning mortality, reduced feed conversion efficiency, increased drug and labor cost and have been estimated to cost approximately \$236 per sow in addition to loss of profits (Polson et al., Financial implications of mystery swine disease (MSD), Proc. Mystery Swine Disease Committee Meeting, Denver, Co., 1990, pp. 8-28). This represents a loss of \$23,600 for a 100 sow herd or \$236,000 for a 1000 sow herd.

presson the exact economic losses from preumonia in nursery pigs. However, the exact economic losses from pressociated preumonia are not known. Presson is an important cause of preumonia in nursery and weared pigs. Reproductive disease was the predominant clinical outcome of presson infections during the past few years. Respiratory disease has now become the main problem associated with presson.

Surprisingly, the market for animal vaccines in the U.S. and worldwide is larger than the market for human vaccines. Thus, there exists an economic incentive to develop new veterinary vaccines, in addition to the substantial public health benefit which is derived from protecting farm animals from disease.

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It is a further object of the present invention to provide an isolated polynucleic acid which encodes a PRRSV protein.

It is a further object of the present invention to provide a PRRSV protein, either isolated from a PRRSV or encoded by a PRRSV polynucleic acid.

It is a further object of the present invention to provide a protein- or polynucleic acid-based vaccine which protects a pig against PRRS.

It is a further object of the present invention to provide a method of raising an effective immunological response against a PRRSV using the vaccine.

It is a further object of the present invention to provide a method of producing a protein- or polynucleic acid-based vaccine which protects a pig against a PRRSV infection.

It is a further object of the present invention to provide a method of treating a pig infected by or exposed to a PRRSV.

It is a further object of the present invention to provide a method of detecting PRRSV.

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It is a further object of the present invention to provide an immunoperoxidase diagnostic assay for detection of PRRSV antigen in porcine tissues.

It is a further object of the present invention to provide an antibody which immunologically binds to a PRRSV protein or to an antigenic region of such a protein.

It is a further object of the present invention to provide an antibody which immunologically binds to a protein- or polynucleic acid-based vaccine which protects a pig against a PRRSV.

It is a further object of the present invention to provide a method of treating a pig exposed to or infected by a PRRSV.

It is a further object of the present invention to provide a method of detecting and a diagnostic kit for assaying a PRRSV.

It is a further object of the present invention to provide the above objects, where the PRRS virus is the Iowa strain of PRRSV.

These and other objects which will become apparent during the following description of the preferred embodiments, have been provided by at least one purified polypeptide selected from the group consisting of proteins encoded by one or more open reading frames (ORF's) of an Iowa strain of porcine reproductive and respiratory virus (PRRSV), proteins at least 80% but less than 100%

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homologous with those encoded by one or more of ORF 2, ORF 3, ORF 4 and ORF 5 of an Iowa strain of PRRSV, proteins at least 97% but less than 100% homologous with proteins encoded by one or both of ORF 6 and ORF 7 of an Iowa strain of PRRSV, antigenic regions of said proteins which are at least 5 amino acids in length and which effectively stimulate immunological protection in a porcine host against a subsequent challenge with a PRRSV isolate, and combinations thereof; an isolated polynucleic acid which encodes such a polypeptide or polypeptides; a vaccine comprising an effective amount of such a polynucleotide or polypeptide(s); antibodies which specifically bind to such a polynucleotide or polypeptide; methods of producing the same; and methods of raising an effective immunological response against a PRRSV, treating a pig exposed to or infected by a PRRSV, and detecting a PRRSV using the same.

## BRIEF DESCRIPTION OF THE FIGURES

Figure 1 is a flowchart outlining a procedure for producing a subunit vaccine;

Figure 2 is a flowchart outlining a procedure for producing a genetically engineered vaccine;

Figure 3 shows a general schematic procedure for the construction of a cDNA  $\lambda$  library as described by the manufacturer (Stratagene)

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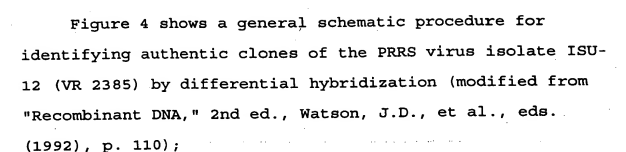


Figure 5 is a Northern blot showing the VR 2385 subgenomic mRNA species, denatured with 6 M glyoxal and DMSO, and separated on a 1.5% agarose gel;

Figure 6 shows the  $\lambda$  cDNA clones used to obtain the 3'-terminal nucleotide sequence of VR 2385;

Figure 7 shows the 2062 bp 3'-terminal sequence (SEC

ID NO:13) and the amino acid sequences encoded by OFF

6 and 7 (SEQ ID NOS:15, 17 and 19, respectively) of VR

2385;

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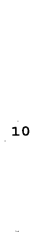
Figure 8 compares the ORF-5 regions of the genomes of VR 2385 and Lelystad virus;

Figure 9 compares the ORF-6 regions of the genomes of VR 2385 and Lelystad virus;

Figure 10 compares the ORF-7 regions of the genomes of VR 2385 and Lelystad virus;

Figure 11 compares the 3'-nontranslational regions of the genomes of VR 2385 and Lelystad virus;

Figure 12 shows a cytopathic effect in HI-FIVE cells infected with a recombinant baculovirus containing the VR 2385 ORF-7 gene (Baculo.PRRSV.7);



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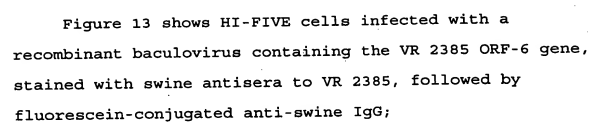


Figure 14 shows HI-FIVE cells infected with a recombinant baculovirus containing the VR 2385 ORF-7 gene, respectively, stained with swine antisera to VR 2385, followed by fluorescein-conjugated anti-swine IgG;

Figure 15 shows a band of expected size for the VR 2385 ORF-6 product, detected by a radioimmunoprecipitation technique (see Experiment II(B) below);

Figure 16 shows a band of expected size for the VR 2385 ORF-7 product, detected by a radioimmunoprecipitation technique (see Experiment II(B) below);

Rigure 17 compares the QRF 6 and QRF / nucleotide sequences of six U.S. PRRSV isolates and of LV, in which the VR 2385 nucleotide sequence is shown first, and in subsequent sequences, only those nucleotides which are different are indicated;

Figures 18(A)-(B) show the alignment of amino acid sequences of the putative M (Fig. 18(A)) and N (Fig. 18(B)) genes of the proposed arterivirus group, performed with a GENEWORKS program (IntelliGenetics, Inc.);

Figures 19(A)-(B) show phylogenetic trees based on the amino acid sequences of the putative M (Fig. 19(A)) and N genes (Fig. 19(B)) for the proposed arterivirus group;

Figures 21(A)-(C) compare the nucleotide sequences of ORF 2, ORF 3 and ORF 4 of PRRSV VR 2385 with the corresponding ORF's of Lelystad virus (LV);

Figures 22(A)-(C) show alignments of the predicted amino acid sequences encoded by ORF's 2, 3 and 4 of PRRSV VR 2385 and LV;

Figure 23 shows an immunohistochemical stain of a lung tissue sample taken from a pig infected 9 days previously with PRRSV, in which positive ABC staining with hematoxylin counterstain is observed within the cytoplasm of macrophages and sloughed cells in the alveolar spaces;

Figure 24 shows an immunohistochemical stain of a lung tissue sample taken from a pig infected 4 days previously with PRRSV, in which positive ABC staining with hematoxylin counterstain is demonstrated within cellular debris in terminal airway lumina;

Figure 25 shows a heart from a pig infected 9 days previously with PRRSV, in which positive staining is demonstrated within endothelial cells (arrow) and isolated macrophages by the present streptavidin-biotin complex method (with hematoxylin counterstain); the bar indicates a length of 21 microns;

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Figure 26 shows a tonsil from a pig infected 9 days previously with PRRSV, in which positive staining cells (arrow heads) are demonstrated within follicles and in the crypt epithelium by the present streptavidin-biotin complex method (with hematoxylin counterstain); the bar indicates a length of 86 microns;

Figure 27 shows a lymph node from a pig infected 9 days previously with PRRSV, in which positive staining is demonstrated within follicles by the present streptavidin-biotin complex method (with hematoxylin counterstain), and positive cells (arrows) resemble macrophages or dendritic cells; the bar indicates a length of 21 microns;

Figures 28(A)-(C) are photomicrographs of lungs from pig inoculated with (A) culture fluid from an uninfected cell line, (B) culture fluid from a cell line infected with a low virulence PRRSV isolate (the lungs show PRRS-A type lesions), and (C) culture fluid from a cell line infected with a high virulence PRRSV isolate (the lungs show PRRS-B type lesions);

Figures 29(A)-(B) illustrate immunohistochemical staining with anti-PRRSV monoclonal antibody of a lung from a pig infected 9 days previously with PRRSV; and

Figures 30(A)-(B) show Northern blots of PRRSV isolates VR 2385pp (designated as "12"), VR 2429 (ISU-22, designated as "22"), VR 2430, designated as "55"), ISU-79

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(designated as "79"), ISU-1894 $^{/}$  (designated as "1894"), and VR 2431, designated as "3927").

# DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENTS

In the present invention, a "porcine reproductive and respiratory syndrome virus" or "PRRSV" refers to a virus which causes the diseases PRRS, PEARS, SIRS, MSD and/or PIP (the term "PIP" now appears to be disfavored), including the Iowa strain of PRRSV, other strains of PRRSV found in the United States (e.g., VR 2332), strains of PRRSV found in Canada (e.g., IAF-exp91), strains of PRRSV found in Europe (e.g., Lelystad virus, PRRSV-10), and closely-related variants of these viruses which may have appeared and which will appear in the future.

The present vaccine is effective if it protects a pig against infection by a porcine reproductive and respiratory syndrome virus (PRRSV). A vaccine protects a pig against infection by a PRRSV if, after administration of the vaccine to one or more unaffected pigs, a subsequent challenge with a biologically pure virus isolate (e.g., VR 2385, VR 2386, or other virus isolate described below) results in a lessened severity of any gross or histopathological changes (e.g., lesions in the lung) and/or of symptoms of the disease, as compared to those changes or symptoms typically caused by the isolate in similar pigs which are unprotected (i.e., relative to an

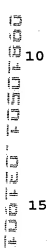
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appropriate control). More particularly, the present vaccine may be shown to be effective by administering the vaccine to one or more suitable pigs in need thereof, then after an appropriate length of time (e.g., 1-4 weeks), challenging with a large sample  $(10^{3-7}\ \text{TCID}_{50})$  of a biologically pure PRRSV isolate. A blood sample is then drawn from the challenged pig after about one week, and an attempt to isolate the virus from the blood sample is then performed (e.g., see the virus isolation procedure exemplified in Experiment VIII below). Isolation of the virus is an indication that the vaccine may not be effective, and failure to isolate the virus is an indication that the vaccine may be effective.

Thus, the effectiveness of the present vaccine may be evaluated quantitatively (i.e., a decrease in the percentage of consolidated lung tissue as compared to an appropriate control group) or qualitatively (e.g., isolation of PRRSV from blood, detection of PRRSV antigen in a lung, tonsil or lymph node tissue sample by an immunoperoxidase assay method [described below], etc.). The symptoms of the porcine reproductive and respiratory disease may be evaluated quantitatively (e.g., temperature/ fever), semi-quantitatively (e.g., severity of respiratory distress [explained in detail below], or qualitatively (e.g., the presence or absence of one or more symptoms or a



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reduction in severity of one or more symptoms, such as cyanosis, pneumonia, heart and/or brain lesions, etc.).

An unaffected pig is a pig which has either not been exposed to a porcine reproductive and respiratory disease infectious agent, or which has been exposed to a porcine reproductive and respiratory disease infectious agent but is not showing symptoms of the disease. An affected pig is one which shows symptoms of PRRS or from which PRRSV can be isolated.

The clinical signs or symptoms of PRRS may include lethargy, respiratory distress, "thumping" (forced expiration), fevers, roughened haircoats, sneezing, coughing, eye edema and occasionally conjunctivitis.

Lesions may include gross and/or microscopic lung lesions, myocarditis, lymphadenitis, encephalitis and rhinitis. The infectious agent may be a single virus, or may be combined with one or more additional infectious agents (e.g., other viruses or bacteria). In addition, less virulent and non-virulent forms of the PRRSV and of Iowa strain have been found, which may cause either a subset of the above symptoms or no symptoms at all. Less virulent and non-virulent forms of PRRSV can be used according to the present invention to provide protection against porcine reproductive and respiratory diseases nonetheless.

Histological lesions in the various porcine diseases are different. Table I below compares physiological

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observations and pathology of the lesions associated with a number of diseases caused by porcine viruses:

TABLE I

Swine Viral Pneumonia Comparative Pathology

Lesion	PRRS(p)	PRRS(0)	SIV	PNP	PRCV	PPMV	Iowa
Type II	+	+++	+	+++	++	++	++++
Inter. thickening	++++	+	+	+	++	++	+
Alveolar exudate	+	+++	++	++	++	++	+++
Airway necrosis	-	•	++++,	++++	+++	+	-
Syncytia	-	++	+/-	++	+	+.*	+++
Encephalitis	+	+++	-	•	•	++	+
Myocarditis	+/-	++	-	•	•	•	+++

wherein "PRRS(p)" represents the published pathology of the PRRS virus, "PRRS(o)" represents the pathology of PRRS virus observed by the present Inventors, "SIV" represents swine influenza A virus, "PRCV" represents porcine respiratory coronavirus, "PPMV" represents porcine paramyxovirus, "Iowa" refers to the strain of PRRSV discovered by the present Inventors, "Type II" refers to Type II pneumocytes (which proliferate in infected pigs), "Inter." refers to interstitial septal infiltration by mononuclear cells, "Airway necrosis" refers to necrosis in terminal airways, and the symbols (-) and (+) through (++++) refer to a comparative severity scale as follows:

(-): negative (not observed)



(++): moderate

(+++): severe

5 (++++): most severe

A "porcine reproductive and respiratory virus" or "PRRSV" causes a porcine reproductive and respiratory disease defined by one or more of the clinical signs, symptoms, lesions and histopathology as described above, and is characterized as being an enveloped RNA arterivirus, having a size of from 50 to 80 nm in diameter and from 250 to 400 nm in length. "North American strains of PRRSV" refer to those strains of PRRSV which are native to North "U.S. strains of PRRSV" refer to strains of PRRSV native to the U.S., and "European strains of PRRSV" refer to strains native to Europe, such as Lelystad virus (deposited by the CDI [Lelystad, Netherlands] in the depository at the Institut Pasteur, Paris, France, under the deposit number I-1102; see International Patent Publication No. WO 92/21375, published on December 10, 1992).

The "Iowa strain" of PRRSV refers to (a) those strains of PRRSV isolated by the presented Inventors, (b) those strains having at least a 97% sequence identity (or homology) in the seventh open reading frame (ORF 7) with at least one of VR 2385, VR 2430 and VR 2431; (c) strains

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which, after no more than 5 passages, grow to a titer of at least 10<sup>4</sup> TCID<sub>50</sub> in CRL 11171 cells, MA-104 cells or PSP-36 cells, (d) those strains having at least 80% and preferably at least 90% homology with one or more of ORF's 2-5 of VR 2385, and (e) those strains which cause a greater percentage consolidation of lung tissue than Lelystad virus (e.g., at 10 days post-infection, infected pigs exhibit at least 20% and preferably at least 40% lung consolidation). Preferably, the Iowa strain of PRRSV is characterized by at least two of the above characteristics (a)-(e).

The present invention is primarily concerned with polynucleic acids (segments of genomic RNA and/or DNA, mRNA, cDNA, etc.) isolated from or corresponding to a porcine reproductive and respiratory syndrome virus (PRRSV), proteins encoded by the DNA, methods of producing the polynucleic acids and proteins, vaccines which protect pigs from a PRRSV, a method of protecting a pig from a PRRSV using the vaccine, a method of producing the vaccine, a method of treating a pig infected by or exposed to a PRRSV, and a method of detecting a PRRSV. particularly, the present invention is concerned with a vaccine which protects pigs from North American strains of PRRSV, a method of producing and administering the vaccine, and polynucleic acids and proteins obtained from an Iowa strain of PRRSV. However, it is believed that the information learned in the course of developing the present



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invention will be useful in developing vaccines and methods of protecting pigs against any and/or all strains of porcine reproductive and respiratory syndrome. Therefore, the present invention is not necessarily limited to polynucleic acids, proteins, vaccines and methods related to the Iowa strain of PRRS virus (PRRSV).

The phrase "polynucleic acid" refers to RNA or DNA, as well as mRNA and cDNA corresponding to or complementary to the RNA or DNA isolated from the virus or infectious agent. An "ORF" refers to an open reading frame, or polypeptide-encoding segment, isolated from a viral genome, including the PRRSV genome. In the present polynucleic acid, an ORF can be included in part (as a fragment) or in whole, and can overlap with the 5'- or 3'-sequence of an adjacent ORF (see Figs. 7 and 21, and Experiments I and IV below). A "polynucleotide" is equivalent to a polynucleic acid, but may define a distinct molecule or group of molecules (e.g., as a subset of a group of polynucleic acids).

Referring now to Figures 1-2, flowcharts of procedures are provided for preparing types of vaccines encompassed by the present invention. The flowcharts of Figures 1-2 are provided as exemplary methods of producing the present vaccines, and are not intended to limit the present invention in any manner.

The first step in each procedure detailed in Figures
1-2 is to identify a cell line susceptible to infection

with a porcine reproductive and respiratory virus or infectious agent. (To simplify the discussion concerning preparation of the vaccine, the term "virus" refers to a virus and/or other infectious agent associated with a porcine reproductive and respiratory disease.) A master cell stock (MCS) of the susceptible host cell is then The susceptible host cells continue to be prepared. passaged beyond MCS. Working cell stock (WCS) is prepared from cell passages between MCS and MCS+n.

A master seed virus is propagated on the susceptible host cell line, between MCS and MCS+n, preferably on WCS. The raw virus is isolated by methods known in the art from appropriate, preferably homogenized, tissue samples taken from infected pigs exhibiting disease symptoms corresponding to those caused by the virus of interest. suitable host cell, preferably a sample of the WCS, is infected with the raw virus, then cultured. Vaccine virus is subsequently isolated and plaque-purified from the infected, cultured host cell by methods known in the art. Preferably, the virus to be used to prepare the vaccine is plaque-purified three times.

Master seed virus (MSV) is then prepared from the plaque-purified virus by methods known in the art. MSV(X) is then passaged in WCS at least four times through MSV(X+1), MSV(X+2), MSV(X+3) and MSV(X+4) virus passages. The MSV(X+4) is considered to be the working seed virus.

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Preferably, the virus passage to be used in the pig studies and vaccine product of the present invention is MSV(X+5), the product of the fifth passage.

In conjunction with the working cell stock, the working seed virus is cultured by known methods in sufficient amounts to prepare a prototype vaccine, preferably MSV(X+5). The present prototype vaccines may be of any type suitable for use in the veterinary medicine field. The primary types of vaccines on which the present invention focuses include a subunit vaccine (Figure 1) and a genetically engineered vaccine (Figure 2). However, other types of vaccines recognized in the field of veterinary vaccines, including live, modified live, attenuated and killed virus vaccines, are also acceptable. A killed vaccine may be rendered inactive through chemical treatment or heat, etc., in a manner known to the artisan of ordinary skill.

An attenuated virus may be obtained by repeating serial passage of the virus in a suitable host cell a sufficient number of times to obtain an essentially non-virulent virus. For example, a PRRSV may be serially passaged from 1 to 20 times (or more, if desired), in order to render it sufficiently attenuated for use as an attenuated vaccine. MSV(X+5) may be such an attenuated vaccine.

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In the procedures outlined by each of Figures 1-2, following preparation of a prototype vaccine, pig challenge models and clinical assays are conducted by methods known in the art. For example, before performing actual vaccination/challenge studies, the disease to be prevented and/or treated must be defined in terms of its symptoms, clinical assay results, conditions, etc. As described herein, the Iowa strain of PRRSV has been defined in terms of its histopathology and the clinical symptoms which it causes. Clinical analyses of the Iowa strain of PRRSV are described in detail in the Experiments below.

One then administers a prototype vaccine to a pig, then exposes the pig to the virus which causes the disease. This is known as "challenging" the pig and its immunological system. After observing the response of the challenged pig to exposure to the virus or infectious agent and analyzing the ability of the prototype vaccine to protect the pig, efficacy studies are then performed by conventional, known methods. A potency assay is then developed in a separate procedure by methods known in the art, and prelicensing serials are then produced.

Prior to preparation of the prototype subunit vaccine (Figure 1), the protective or antigenic components of the vaccine virus should be identified. Such protective or antigenic components include certain amino acid segments or fragments of the viral proteins (preferably coat proteins)

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which raise a particularly strong protective or immunological response in pigs; such antigenic protein fragments fused to non-PRRSV proteins which act as a carrier and/or adjuvant; single or multiple viral coat proteins themselves, oligomers thereof, and higher-order associations of the viral coat proteins which form virus substructures or identifiable parts or units of such substructures; oligoglycosides, glycolipids or glycoproteins present on or near the surface of the virus or in viral substructures such as the nucleocapsid; lipoproteins or lipid groups associated with the virus, etc.

Antigenic amino acid segments or fragments are preferably at least 5 amino acids in length, particularly preferably at least 10 amino acids in length, and can be up to but not including the entire length of the native protein. In the present invention, the binding affinity (or binding constant or association constant) of an antigenic fragment is preferably at least 1% and more preferably at least 10% of the binding affinity of the corresponding full-length protein (i.e., which is encoded by the same ORF) to a monoclonal antibody which specifically binds the full-length protein. The monoclonal antibody which specifically binds to the full-length protein encoded by an ORF of a PRRSV is preferably deposited under the Budapest Treaty at an acceptable

depository, or is sequenced or otherwise characterized in terms of its physicochemical properties (e.g., antibody type [IgG, IgM, etc.], molecular weight, number of heavy and light chains, binding affinities to one or more known or sequenced proteins [e.g., selected from SEQ ID NOS:15, 17, 19, 21, 24, 26, 43, 45, 47, 49, 51, 53, 55, 57, 59, 61, 67, 69, 71, 73, 75 and 77], etc.).

Antigenic fragments of viral proteins (e.g., those encoded by one or more of ORF's 2-6 of a PRRSV virus) are identified by methods known in the art. For example, one can prepare polynucleic acids having a truncated ORF encoding a polypeptide with a predetermined number of amino acid residues deleted from the N-terminus, C-terminus, or The truncated ORF can be expressed in vitro or in vivo in accordance with known methods, and the corresponding truncated polypeptide can then be isolated in The immunoprotective accordance with known methods. properties of the polypeptides may be measured directly (e.g., in vivo). Alternatively, the antigenic region(s) of the full-length polypeptide can be determined indirectly by screening a series of truncated polypeptides against, for example, suitably deposited or characterized monoclonal (If the alternative, indirect method is antibodies. performed, the failure of a truncated polypeptide to bind to a neutralizing monoclonal antibody is a strong indication that the portion of the full-length polypeptide

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deleted in the truncated polypeptide contains an antigenic fragment.) Once identified, the antigenic or immunoprotective portion(s) (the "subunit(s)") of the viral proteins or of the virus itself may be subsequently cloned and/or purified in accordance with known methods. (The viral/bacterial inactivation and subunit purification protocols recited in Fig. 1 are optional.)

Genetically engineered vaccines (Figure 2) begin with a modification of the general procedure used for preparation of the other vaccines. After plaque-purification, the PRRS virus may be isolated from a suitable tissue homogenate by methods known in the art, preferably by conventional cell culture methods using PSP-36, ATCC CRL 11171 or macrophage cells as hosts.

The RNA is extracted from the biologically pure virus by a known method, preferably by the guanidine isothiocyanate method using a commercially available RNA isolation kit (for example, the kit available from Stratagene, La Jolla, California), and purified by one or more known methods, preferably by ultracentrifugation in a CsCl gradient. Messenger RNA may be further purified or enriched by oligo (dT)-cellulose column chromatography.

The viral genome is then cloned into a suitable host by methods known in the art (see <u>Maniatis et al</u>, "Molecular Cloning: A Laboratory Manual," Cold Spring Harbor Laboratory (1989), Cold Spring Harbor, Massachusetts). The

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virus genome is then analyzed to determine essential regions of the genome for producing antigenic portions of the virus. Thereafter, the procedure for producing a genetically engineered vaccine is essentially the same as for a modified live vaccine, an inactivated vaccine or a subunit vaccine (see Figure 1 of the present application and Figures 1-3 of U.S. application Serial No. 08/131,625). During prelicensing serials, expression of the cloned, recombinant subunit of a subunit vaccine may be optimized by methods known to those in the art (see, for example, relevant sections of Maniatis et al, cited above).

The present vaccine protects pigs against a virus or infectious agent which causes a porcine reproductive and respiratory disease. Preferably, the present vaccine protects pigs against infection by PRRSV. However, the present vaccine is also expected to protect a pig against infection by closely related variants of various strains of PRRSV as well.

Subunit virus vaccines may also be prepared from semipurified virus subunits by the methods described above in
the discussion of Figure 1. For example, hemagglutinin
isolated from influenza virus and neuraminidase surface
antigens isolated from influenza virus have been prepared,
and shown to be less toxic than the whole virus. Subunit
vaccines can also be prepared from highly purified subunits
of the virus. An example in humans is the 22-nm surface

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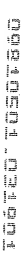
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antigen of human hepatitis B//virus. Human herpes simplex virus subunits and many other examples of subunit vaccines for use in humans are known. Thus, methods of preparing purified subunit vaccines from PRRSV cultured in a suitable host cell may be applicable to the present subunit vaccine.

Attenuated virus vaccines can be found in nature and may have naturally-occurring gene deletions (see Experiments VIII and IX below). Alternatively, attenuated vaccines may be prepared by a variety of known methods, such as serial passage (e.g., 5-25 times) in cell cultures or tissue cultures. However, the attenuated virus vaccines preferred in the present invention are those attenuated by recombinant gene deletions or gene mutations (as described above).

Genetically engineered vaccines are produced by techniques known to those in the art. Such techniques include those using recombinant DNA and those using live viruses. For example, certain virus genes can be identified which code for proteins responsible for inducing a stronger immune or protective response in pigs. Such identified genes can be cloned into protein expression vectors, such (but not limited to) as the baculovirus vector (see, for example, O'Reilly et al, "Baculovirus Expression Vectors: A Lab Manual, "Freeman & Co. (1992)). The expression vector containing the gene encoding the immunogenic virus protein can be used to infect appropriate



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host cells. The host cells are cultured, thus expressing the desired vaccine proteins, which can be purified to a desired extent, then used to protect the pigs from a reproductive and respiratory disease.

Genetically engineered proteins may be expressed, for example, in insect cells, yeast cells or mammalian cells. The genetically engineered proteins, which may be purified and/or isolated by conventional methods, can be directly inoculated into animals to confer protection against porcine reproductive and respiratory diseases. One or more envelope proteins from a PRRSV (i.e., those encoded by ORF's 2-6) or antigenic portions thereof may be used in a vaccine to induce neutralizing antibodies. Nucleoproteins from a PRRSV may be used in a vaccine to induce cellular immunity.

Preferably, the present invention transforms an insect cell line (HI-FIVE) with a transfer vector containing polynucleic acids obtained from the Iowa strain of PRRSV. Preferably, the present transfer vector comprises linearized baculovirus DNA and a plasmid containing one or more polynucleic acids obtained from the Iowa strain of PRRSV. The host cell line may be co-transfected with the linearized baculovirus DNA and a plasmid, so that a recombinant baculovirus is made. Particularly preferably, the present polynucleic acid encodes one or more proteins of the Iowa strain of PRRSV.





Alternatively, RNA or DNA from a PRRSV encoding one or more viral proteins (e.g., envelope and/or nucleoproteins) can be inserted into live vectors, such as a poxvirus or an adenovirus, and used as a vaccine.

Thus, the present invention further concerns a purified preparation of a polynucleic acid isolated from the genome of a PRRS virus, preferably a polynucleic acid isolated from the genome of the Iowa strain of PRRSV. The present polynucleic acid has utility (or usefulness) in the production of the present vaccine, in screening or identifying infected or exposed animals, in identifying related viruses and/or infectious agents, and as a vector for transforming cells and/or immunizing animals (e.g., pigs) with heterologous genes.

In the Experiments described hereinbelow, the isolation cloning and sequencing of ORF's 2-7 of plaque-purified PRRSV isolate ISU-12 (deposited on October 30, 1992, in the American Type Culture Collection, 12301

Parklawn Drive, Rockville, Maryland 20852, U.S.A. under the accession numbers VR 2385 [3 x plaque-purified] and VR 2386 [non-plaque-purified]) and ORF's 6-7 of PRRSV isolates ISU-22, ISU-55 and ISU-3927 (deposited on September 29, 1993, in the American Type Culture Collection under the accession numbers VR 2429, WR 2430 and VR 2431, respectively) ISU-79 and ISU-1894 (deposited on August 31,

respectively) ISU-79 and ISU-1894 (deposited on August 31, 1994, in the American Type Culture Collection under the

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described in detail. However, the techniques used to isolate, clone and sequence these genes can be also applied to the isolation, cloning and sequencing of the genomic polynucleic acids of any PRRSV. Thus, the present invention is not limited to the specific sequences disclosed in the Experiments below.

For example, primers for making relatively large amounts of DNA by the polymerase chain reaction (and if desired, for making RNA by transcription and/or protein by translation in accordance with known in vivo or in vitro methods) can be designed on the basis of sequence information where more than one sequence obtained from a PRRSV genome has been determined (e.g., ORF's 2-5 of VR 2385 and Lelystad virus, or ORF's 6-7 of VR 2385, VR 2429, VR 2430, ISU-79, ISU-1894, VR 2431 and Lelystad virus). A region from about 15 to 50 nucleotides in length having at least 80% and preferably at least 90% identity/is selected ) from the determined sequences. A region where a deletion occurs in one of the sequences (e.g., of at least 5 nucleotides) can be used as the basis for preparing a selective primer for selective amplification of the polynucleic acid of one strain or type of PRRSV over another (e.g., for the differential diagnosis of North American and European PRRSV strains).

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Once the genomic polynucleic acid is amplified and cloned into a suitable host by known methods, the clones can be screened with a probe designed on the basis of the sequence information disclosed herein. For example, a region of from about 50 to about 500 nucleotides in length is selected on the basis of either a high degree of identity (e.g., at least 90%) among two or more sequences (e.g., in ORF's 6-7 of the Iowa strains of PRRSV disclosed in Experiment III below), and a polynucleotide of suitable length and sequence identity can be prepared by known methods (such as automated synthesis, or restriction of a suitable fragment from a polynucleic acid containing the selected region, PCR amplification using primers which hybridize specifically to the polynucleotide, and isolation by electrophoresis). The polynucleotide may be labeled with, for example, 32P (for radiometric identification) or biotin (for detection by fluorometry). The probe is then hybridized with the polynucleic acids of the clones and detected according to known methods.

The present Inventors have discovered that ORF 4 appears to be related to the virulence of PRRSV. For example, at least one isolate of PRRSV which shows relatively low virulence also appears to have a deletion in ORF 4 (see, for example, Experiments VIII-XI below). Accordingly, in a preferred embodiment, the present

invention is concerned with a polynucleic acid obtained

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from a PRRSV isolate which confers immunogenic protection directly or indirectly against a subsequent challenge with a PRRSV, but in which ORF 4 is deleted or mutated to an extent which would render a PRRSV containing the polynucleic acid either low-virulent (i.e., a "low virulence" (lv) phenotype; see the explanation below) or non-virulent (a so-called "deletion mutant"). Preferably, ORF 4 is deleted or mutated to an extent which would render a PRRS virus non-virulent. However, it may be desirable to retain regions of a PRRSV ORF 4 in the present polynucleic acid which (i) encode an antigenic, immunoprotective peptide fragment and (ii) would not confer virulence to a PRRS virus containing the polynucleic acid.

The present invention also encompasses a PRRSV per se in which ORF 4 is deleted or mutated to an extent which renders it either low-virulent or non-virulent (e.g., VR 2431). Such a virus is useful as a vaccine or as a vector for transforming a suitable host (e.g., MA-104, PSP 36, CRL 11171, MARC-145 or porcine alveolar macrophage cells) with a heterologous gene. Preferred heterologous genes which may be expressed using the present deletion mutant may include those encoding a protein or an antigen other than a porcine reproductive and respiratory syndrome virus antigen (e.g., pseudorabies and/or swine influenza virus proteins and/or polypeptide-containing antigens, a porcine growth



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hormone, etc.) or a polypeptide-based adjuvant (such as those discussed below for the present vaccine composition).

It may also be desirable in certain embodiments of the present polynucleic acid which contain, for example, the 3'-terminal region of ORF 3 (e.g., from 200 to 700 nucleotides in length), at least part of which may overlap with the 5'-region of ORF 4. Similarly, where the 3'terminal region of ORF 4 may overlap with the 5'-terminal region of ORF 5, it may be desirable to retain the 5'region of ORF 4 which overlaps with ORF 5.

The present Inventors have also discovered that ORF 5 in the PRRSV genome appears to be related to replication of the virus in mammalian host cells capable of sustaining a culture while infected with PRRSV. Accordingly, the present invention is also concerned with polynucleic acids obtained from a PRRSV genome in which ORF 5 may be present in multiple copies (a so-called "overproduction mutant"). For example, the present polynucleic acid may contain at least two, and more preferably, from 2 to 10 copies of ORF 5 from a high-replication (hr) phenotype PRRSV isolate.

Interestingly, the PRRSV isolate ISU-12 has a surprisingly large number of potential start codons (ATG/AUG sequences) near the 5'-terminus of ORF 5, possibly indicating alternate start sites of this gene (see SEQ ID Thus, alternate forms of the protein encoded by NO:13): ORF 5 of a PRRSV isolate may exist, particularly where

alternate ORF's encode a protein having a molecular weight similar to that determined experimentally (e.g., from about 150 to about 250 amino acids in length). The most likely coding region for ORF 5 of ISU-12 (SEQ ID NO:14) is indicated in Figure 7.

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One can prepare deletion and overproduction mutants in accordance with known methods. For example, one can prepare a mutant polynucleic acid which contains a "silent" or degenerate change in the sequence of a region encoding a polypeptide. By selecting and making an appropriate degenerate mutation, one can substitute a polynucleic acid sequence recognized by a known restriction enzyme. example, if such a silent, degenerate mutation is made at one or two of the 3'-end of ORF 3 and the 5'- and 3'-ends of ORF 4 and ORF 5, one can insert a synthetic polynucleic acid (a so-called "cassette") which may contain multiple copies of ORF 5, multiple copies of a viral envelope The "cassette" protein or an antigenic fragment thereof. may be preceded by a suitable initiation codon (ATG), and may be suitably terminated with a termination codon at the 3'-end (TAA, TAG or TGA).

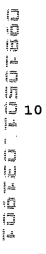
Of course, an oligonucleotide sequence which does not encode a polypeptide may be inserted, or alternatively, no cassette may be inserted. By doing so, one may provide a so-called deletion mutant.

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Thus, in one embodiment of the present invention, the polynucleic acid encodes one or more proteins, or antigenic regions thereof, of a PRRSV. Preferably, the present nucleic acid encodes at least one antigenic region of a PRRSV membrane (envelope) protein. More preferably, the present polynucleic acid contains at least one copy of the ORF-5 gene from a high virulence (hv) phenotype isolate of PRRSV (see the description of "hv phenotype" below) and a sufficiently long fragment, region or sequence of at least one of ORF-2, ORF-3, ORF-4, ORF-5 and/or ORF-6 from the genome of a PRRSV isolate to encode an antigenic region of the corresponding protein(s) and effectively stimulate immunological protection against a subsequent challenge with an hv phenotype PRRSV isolate. Even more preferably, at least one entire envelope protein encoded by ORF-2, ORF-3, ORF-5 and/or ORF-6 of a PRRSV is contained in the present polynucleic acid, and the present polynucleic acid excludes a sufficiently long portion of ORF 4 from an hv PRRSV to render a PRRSV containing the same either lowvirulent or non-virulent. Particularly preferably, the present polynucleic acid excludes the entire region of an hv PRRSV ORF 4 which does not overlap with the 3'-end of ORF 3 and the 5'-end of ORF 5.

Most preferably, the polynucleic acid is isolated from the genome of an isolate of the Iowa strain of PRRSV (for example, VR 2385 (3X plaque-purified ISU-12), VR 2386 (non-



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plaque-purified ISU-12), VR 2428 (ISU-51), VR 2429 (ISU-22), VR 2430 (ISU-55), VR 2431 (ISU-3927), ISU-79 and/or ISU-1894.

A preferred embodiment of the present invention concerns a purified preparation which may comprise, consist essentially of or consist of a polynucleic acid having a sequence of the formula (I):

$$5'-\alpha-\beta-\gamma-3'$$
 (I)

wherein  $\alpha$  encodes at least one polypeptide or antigenic fragment thereof encoded by a polynucleotide selected from the group consisting of ORF 2 and ORF 3 of an Iowa strain of PRRSV and regions thereof encoding the antigenic fragments; and  $\beta$  is either a covalent bond or a linking polynucleic acid which excludes a sufficiently long portion of ORF 4 from an hv PRRSV to render the hv PRRSV either low-virulent or non-virulent; and  $\gamma$  is at least one copy of an ORF 5 from an Iowa strain of PRRSV, preferably from a high replication (hr) phenotype.

Alternatively, the present invention may concern a purified preparation which may comprise, consist essentially of or consist of a polynucleic acid having a sequence of the formula (II):

(II)

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where  $\gamma$  is at least one copy of an ORF 5 from an Iowa strain of PRRSV, preferably from an hv PRRSV isolate;  $\delta$  is either a covalent bond or a linking polynucleic acid which does not materially affect transcription and/or translation of the polynucleic acid; and  $\epsilon$  encodes at least one polypeptide or antigenic fragment thereof encoded by a polynucleotide selected from the group consisting of ORF 6 and ORF 7 of an Iowa strain of PRRSV and regions thereof encoding the antigenic fragments; and when  $\delta$  is a covalent bond,  $\gamma$  may have a 3'-end which excludes the region overlapping with the 5'-end of a corresponding ORF 6. Preferably,  $\epsilon$  is a polynucleotide encoding at least an antigenic region of a protein encoded by an ORF 6 of an Iowa strain of PRRSV, and more preferably, encodes at least a protein encoded by an ORF 6 of an Iowa strain of PRRSV.

The present invention may also concern a purified preparation which may comprise, consist essentially of or consist of a polynucleic acid having a sequence of the formula (III):

 $5'-\alpha-\beta-\gamma-\delta-\epsilon-3'$  (III)

where  $\alpha$ ,  $\beta$ ,  $\gamma$ ,  $\delta$  and  $\epsilon$  are as defined in formulas (I) and (II) above. Thus, the present polynucleic acid may be selected from the group consisting of, from 5' to 3':

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(ORF 5) <sub>n</sub>	(IV)
$\zeta$ -(ORF 5) <sub>n</sub>	(v)
$(ORF 5)_n - \eta$	(VI)
ζ-(ORF 5) <sub>n</sub> -η	(VII)

#### 5 where:

 $\eta$  is selected from the group consisting of -ORF 5\*, -ORF 6, -ORF 7, -ORF 5\*-ORF 6, -ORF 5\*-ORF 7 and -ORF 5\*-ORF 6-ORF 7;

wherein ORF 2, ORF 3, ORF 6 and ORF 7 each encode a protein encoded by the second, third, sixth and seventh open reading frames of an Iowa strain of PRRSV, respectively; ORF 4\* is a region of a fourth open reading frame of an Iowa strain of PRRSV which (i) encodes an antigenic, immunoprotective peptide fragment and which (ii) does not confer virulence to a PRRSV containing the polynucleic acid; ORF 5 is a fifth open reading frame of an hv PRRSV isolate; ORF 5\* is a region of a fifth open reading frame of an Iowa strain of PRRSV which (i) encodes an antigenic, immunoprotective peptide fragment and (ii) does not confer virulence to a PRRSV containing the polynucleic acid, and which may have a 3'-end which excludes the

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portion overlapping with the 5'-end of a corresponding ORF 6; and  $n \ge 1$ .

The present polynucleic acid may also comprise, consist essentially of or consist of combinations of the above sequences, either as a mixture of polynucleotides or covalently linked in either a head-to-tail (sense-antisense) or head-to-head fashion. Polynucleic acids complementary to the above sequences and combinations thereof (antisense polynucleic acid) are also encompassed by the present invention. Thus, in addition to possessing multiple or variant copies of ORF 5, the present polynucleic acid may also contain multiple or variant copies of one or more of ORF's 1-3 and 6-7 and regions of ORF's 4-5 of Iowa strain PRRSV's.

The present invention may also concern polynucleic acids comprising, consisting essentially of or consisting of the open reading frame 1a and 1b from a PRRSV isolate. Based on information regarding viruses evolutionally related to PRRSV, ORF 1a and 1b of PRRSV are believed to encode an RNA polymerase. ORF 1a and 1b are translated into a single protein by frameshifting. Preferably, the polynucleic acid from ORF 1a and 1b of a PRRSV isolate is obtained from an Iowa strain of PRRSV.

Similar to the methods described above and in the following Experiments for ORF's 2-7, one can prepare a library of recombinant clones (e.g., using *E. coli* as a





host) containing suitably prepared restriction fragments of a PRRSV genome (e.g., inserted into an appropriate plasmid expressible in the host). The clones are then screened with a suitable probe (e.g., based on a conserved sequence of ORF's 2-3; see, for example, Figure 22). Positive clones can then be selected and grown to an appropriate level. The polynucleic acids can then be isolated from the positive clones in accordance with known methods. A suitable primer for PCR can then be designed and prepared as described above to amplify the desired region of the polynucleic acid. The amplified polynucleic acid can then be isolated and sequenced by known methods.

The present purified preparation may also contain a polynucleic acid selected from the group consisting of sequences having at least 97% sequence identity (or homology) with at least one ORF 7 of VR 2385, VR 2430 and/or VR 2431; and sequences having at least 80% and preferably at least 90% sequence identity (or homology) with at least one of ORF's 1-6 of VR 2385, VR 2428, VR 2429, VR 2430 and/or VR 2431. Preferably, the polynucleic acid excludes a sufficiently long region or portion of ORF 4 of the hv PRRSV isolates VR 2385, VR 2429, ISU-28, ISU-79 and/or ISU-984 to render the isolate low-virulent or non-virulent.

In the context of the present application, "homology" refers to the percentage of identical nucleotide or amino

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acid residues in the sequences of two or more viruses, aligned in accordance with a conventional method for determining homology (e.g., the MACVECTOR or GENEWORKS computer programs, aligned in accordance with the procedure described in Experiment III below).

Accordingly, a further aspect of the present invention encompasses an isolated polynucleic acid at least 90% homologous to a polynucleotide which encodes a protein, polypeptide or fragment thereof encoded by ORF's 1-7 from an Iowa strain of PRRSV (e.g., SEQ ID NOS:15, 17, 19, 43, 45, 47, 49, 51, 53, 55, 57, 59, 61, 63, 65 and 67). Preferably, the present isolated polynucleic acid encodes a protein, polypeptide, or antigenic fragment thereof which is at least 10 amino acids in length and in which amino acids non-essential for antigenicity may be conservatively substituted. An amino acid residue in a protein, polypeptide, or antigenic fragment thereof is conservatively substituted if it is replaced with a member of its polarity group as defined below:

#### Basic amino acids:

lysine (Lys), arginine (Arg), histidine (His)

Acidic amino acids:

aspartic acid (Asp), glutamic acid (Glu), asparagine (Asn), glutamine (Gln)



serine (Ser), threonine (Thr), cysteine (Cys), asparagine (Asn), glutamine (Gln)

# Sulfur-containing amino acids:

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1- 1-1 mg 1.1 1- 1-1 1.2 1.3 1.3

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cysteine (Cys), methionine (Met)

### Hydrophobic, aromatic amino acids:

phenylalanine (Phe), tyrosine (Tyr), tryptophan (Trp)

# Hydrophobic, nonaromatic amino acids:

glycine (Gly), alanine (Ala), valine (Val), leucine (Leu), isoleucine (Ile), proline (Pro)

More particularly, the present polynucleic acid encodes one or more of the protein(s) encoded by the second, third, fourth, fifth, sixth and/or seventh open reading frames (ORF's 2-7) of the PRRSV isolates VR 2385, VR 2386, VR 2428, VR 2429, VR 2430, VR 2431, VR 2432, ISU-79 and/or ISU-1894 (e.g., SEQ ID NOS:15, 17, 19, 43, 45, 47, 49, 51, 53, 55, 57, 59, 61, 63 and 65).

Relatively short segments of polynucleic acid (about 20 bp or longer) in the genome of a virus can be used to screen or identify tissue and/or biological fluid samples from infected animals, and/or to identify related viruses, by methods described herein and known to those of ordinary skill in the fields of veterinary and viral diagnostics and veterinary medicine. Accordingly, a further aspect of the



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present invention encompasses an isolated (and if desired, purified) polynucleic acid consisting essentially of a fragment of from 15 to 2000 bp, preferably from 18 to 1000 bp, and more preferably from 21 to 100 bp in length, derived from ORF's 2-7 of a PRRSV genome (preferably the Iowa strain of PRRSV). Particularly preferably, the present isolated polynucleic acid fragments are obtained from a terminus of one or more of ORF's 2-7 of the genome of the Iowa strain of PRRSV, and most preferably, are selected from the group consisting of SEQ ID NOS:1-12, 22 and 28-34.

The present invention also concerns a diagnostic kit for assaying a porcine reproductive and respiratory syndrome virus, comprising (a) a first primer comprising a polynucleotide having a sequence of from 10 to 50 nucleotides in length which hybridizes to a genomic polynucleic acid from an Iowa strain of porcine reproductive and respiratory syndrome virus at a temperature of from 25 to 75°C, (b) a second primer comprising a polynucleotide having a sequence of from 10 to 50 nucleotides in length, said sequence of said second primer being found in said genomic polynucleic acid from said Iowa strain of porcine reproductive and respiratory syndrome virus and being downstream from the sequence to which the first primer hybridizes, and (c) a reagent which enables detection of an amplified polynucleic acid.



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Preferably, the reagent is an intercalating dye, the fluorescent properties of which change upon intercalation into double-stranded DNA.

ORF's 6 and 7 are not likely candidates for controlling virulence and replication phenotypes of PRRSV, as the nucleotide sequences of these genes are highly conserved among high virulence (hv) and low virulence (lv) isolates (see Experiment III below). However, ORF 5 in PRRSV isolates appears to be less conserved among high replication (hr) and low replication (lr) isolates. Therefore, it is believed that the presence of an ORF 5 from an hr PRRSV isolate in the present polynucleic acid will enhance the production and expression of a recombinant vaccine produced from the polynucleic acid.

Accordingly, it is preferred that the present polynucleic acid, when used for immunoprotective purposes (e.g., in the preparation of a vaccine), contain at least one copy of ORF 5 from a high-replication isolate (i.e., an isolate which grows to a titer of 10<sup>6</sup>-10<sup>7</sup> TCID<sub>50</sub> in, for example, CRL 11171 cells; also see the discussions in Experiments VIII-XI below).

On the other hand, the lv isolate VR 2431 appears to be a deletion mutant, relative to hv isolates (see Experiments III and VIII-XI below). The deletion appears to be in ORF 4, based on Northern blot analysis.

Accordingly, when used for immunoprotective purposes, the

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present polynucleic acid preferably does not contain a region of ORF 4 from an hv isolate responsible for its high virulence, and more preferably, excludes the region of ORF 4 which does not overlap with the adjacent ORF's 3 and 5 (where ORF 4 overlaps with the adjacent ORF's 3 and 5).

It is also known (at least for PRRSV) that neither the nucleocapsid protein nor antibodies thereto confer immunological protection against the virus (e.g., PRRSV) to Accordingly, the present polynucleic acid, when used pigs. for immunoprotective purposes, contains one or more copies of one or more regions from ORF's 2, 3, 4, 5 and 6 of a PRRSV isolate encoding an antigenic region of the viral envelope protein, but which does not result in the symptoms or histopathological changes associated with PRRS. Preferably, this region is immunologically cross-reactive with antibodies to envelope proteins of other PRRSV Similarly, the protein encoded by the present isolates. immunoprotective polynucleic acid confers immunological protection to a pig administered a composition comprising the protein, and antibodies to this protein are immunologically cross-reactive with the envelope proteins of other PRRSV isolates. More preferably, the present immunoprotective polynucleic acid encodes the entire envelope protein of a PRRSV isolate or a protein at least 80% homologous thereto and in which non-homologous residues are conservatively substituted, or a protein at least 90% homologous thereto.

The present isolated polynucleic acid fragments can be obtained by digestion of the cDNA corresponding to (complementary to) the viral polynucleic acids with one or more appropriate restriction enzymes, can be amplified by PCR and cloned, or can be synthesized using a commercially available automated polynucleotide synthesizer.

Another embodiment of the present invention concerns one or more proteins or antigenic fragments thereof from a PRRS virus, preferably from the Iowa strain of PRRSV. As described above, an antigenic fragment of a protein from a PRRS virus (preferably from the Iowa strain of PRRSV) is at least 5 amino acids in length, particularly preferably at least 10 amino acids in length, and provides or stimulates an immunologically protective response in a pig administered a composition containing the antigenic fragment.

Methods of determining the antigenic portion of a protein are known to those of ordinary skill in the art (see the description above). In addition, one may also determine an essential antigenic fragment of a protein by first showing that the full-length protein is antigenic in a host animal (e.g., a pig). If the protein is still antigenic in the presence of an antibody which specifically binds to a particular region or sequence of the protein,

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then that region or sequence may be non-essential for immunoprotection. On the other hand, if the protein is no longer antigenic in the presence of an antibody which specifically binds to a particular region or sequence of the protein, then that region or sequence is considered to be essential for antigenicity.

The present invention also concerns a protein or antigenic fragment thereof encoded by one or more of the polynucleic acids defined above, and preferably by one or more of the ORF's of a PRRSV, more preferably of the Iowa strain of PRRSV. The present proteins and antigenic fragments are useful in immunizing pigs against PRRSV, in serological tests for screening pigs for exposure to or infection by PRRSV (particularly the Iowa strain of PRRSV), etc.

For example, the present protein may be selected from the group consisting of the proteins encoded by ORF's 2-7 of VR 2385, ISU-22 (VR 2429), ISU-55 (VR 2430), ISU-1894, ISU-79 and ISU-3927 (VR 2431) (e.g., SEQ ID NOS:15, 17, 19, 43, 45, 47, 49, 51, 53, 55, 57, 59, 61, 67, 69 and 71); antigenic regions of at least one of the proteins of SEQ ID SEQ ID NOS:15, 17, 19, 43, 45, 47, 49, 51, 53, 55, 57, 59, 61, 67, 69 and 71 having a length of from 5 amino acids to less than the full length of the polypeptides of SEQ ID NOS:15, 17, 19, 43, 45, 47, 49, 51, 53, 55, 57, 59, 61, 67, 69 and 71; polypeptides at least 80% homologous with a

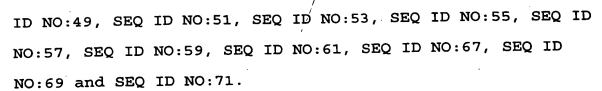
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protein encoded by one of the ORF's 2-5 of VR 2385 (SEQ ID NOS:15, 67, 69 and 71); and polypeptides at least 97% homologous with a protein encoded by one of the ORF's 6-7 of VR 2385, VR 2429, VR 2430, ISU-1894, ISU-79 and VR 2431 (e.g., SEQ ID NOS:17, 19, 43, 45, 47, 49, 51, 53, 55, 57, 59 and 61). Preferably, the present protein has a sequence selected from the group consisting of SEQ ID NOS:15, 17, 19, 43, 45, 47, 49, 51, 53, 55, 57, 59, 61, 67, 69 and 71; variants thereof which provide effective immunological protection to a pig administered the same and in which from 1 to 100 (preferably from 1 to 50 and more preferably from 1 to 25) deletions or conservative substitutions in the amino acid sequence exist; and antigenic fragments thereof at least 5 and preferably at least 10 amino acids in length which provide effective immunological protection to a pig administered the same.

More preferably, the present protein variant or protein fragment has a binding affinity (or association constant) of at least 1% and preferably at least 10% of the binding affinity of the corresponding full-length, naturally-occurring protein to a monoclonal antibody which specifically binds to the full-length, naturally-occurring protein (i.e., the protein encoded by a PRRSV ORF). Most preferably, the present protein has a sequence selected from the group consisting of SEQ ID NO:15, SEQ ID NO:17, SEQ ID NO:19, SEQ ID NO:43, SEQ ID NO:45, SEQ ID NO:47, SEQ



The present invention may also concern a biologically pure virus, characterized in that it contains the present polynucleic acid and/or that it causes a porcine reproductive and respiratory disease which may include one or more of the following histological lesions: gross and/or microscopic lung lesions (e.g., lung consolidation), Type II pneumocytes, myocarditis, encephalitis, alveolar exudate formation and syncytia formation. The phrase "biologically pure" refers to a sample of a virus or infectious agent in which all progeny are derived from a single parent. Usually, a "biologically pure" virus sample is achieved by 3 x plaque purification in cell culture.

In particular, the present biologically pure virus or infectious agent is an isolate of the Iowa strain of pordine reproductive and respiratory syndrome virus, samples of which have been deposited under the terms of the Budapest Treaty at the American Type Culture Collection, 12301 Parklawn Drive, Rockville, Maryland 20852, U.S.A., under the accession numbers VR 2885, VR 2386, VR 2428, VR 2429, VR 2430, VR 2431, 12784 and VALUE.

In addition to the characteristics (a)-(e) described above, the Iowa strain of PRRSV may also be characterized by Northern blots of its mRNA. For example, the Iowa

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strain of PRRSV may contain either 7 or 9 mRNA's, and may also have deletions or variations in their size. In particular, as will be described in the Experiments below, the mRNA's of the Iowa strain of PRRSV may contain up to four deletions, relative to VR 2385/VR 2386.

The present invention further concerns a composition for protecting a pig from viral infection, comprising an amount of the present vaccine effective to raise an immunological response to a virus which causes a porcine reproductive and respiratory disease in a physiologically acceptable carrier.

An effective amount of the present vaccine is one in which a sufficient immunological response to the vaccine is raised to protect a pig exposed to a virus which causes a porcine reproductive and respiratory disease or related illness. Preferably, the pig is protected to an extent in which from one to all of the adverse physiological symptoms or effects (e.g., lung lesions) of the disease to be prevented are found to be significantly reduced.

The composition can be administered in a single dose, or in repeated doses. Dosages may contain, for example, from 1 to 1,000 micrograms of virus-based antigen (vaccine), but should not contain an amount of virus-based antigen sufficient to result in an adverse reaction or physiological symptoms of infection. Methods are known in



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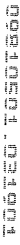
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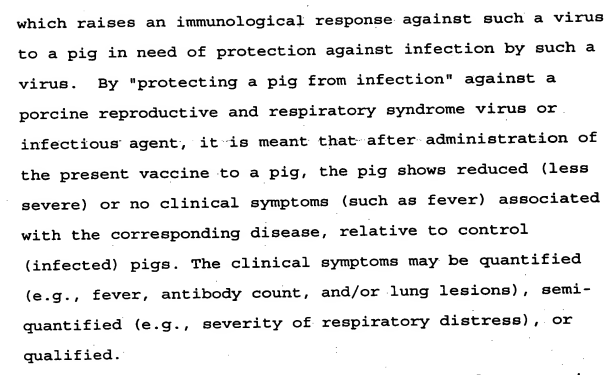
the art for determining suitable dosages of active antigenic agent.

The composition containing the present vaccine may be administered in conjunction with an adjuvant or with an acceptable carrier which may prolong or sustain the immunological response in the host animal. An adjuvant is a substance that increases the immunological response to the present vaccine when combined therewith. The adjuvant may be administered at the same time and at the same site as the vaccine or at a different time, for example, as a booster. Adjuvants also may advantageously be administered to the animal in a manner or at a site or location different from the manner, site or location in which the vaccine is administered. Adjuvants include aluminum hydroxide, aluminum potassium sulfate, heat-labile or heatstable enterotoxin isolated from Escherichia coli, cholera toxin or the B subunit thereof, diphtheria toxin, tetanus toxin, pertussis toxin, Freund's incomplete adjuvant, Freund's complete adjuvant, and the like. Toxin-based adjuvants, such as diphtheria toxin, tetanus toxin and pertussis toxin, may be inactivated prior to use, for example, by treatment with formaldehyde.

The present invention also concerns a method of protecting a pig from infection against a virus which causes a porcine reproductive and respiratory disease, comprising administering an effective amount of a vaccine



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The present invention concerns a system for measuring respiratory distress in affected pigs. The present clinical respiratory scoring system evaluates the respiratory distress of affected pigs by the following scale:

- 0 = no disease; normal breathing
- 1 = mild dyspnea and polypnea when the pigs are stressed (forced to breathe in larger volumes and/or at an accelerated rate)
- 2 = mild dyspnea and polypnea when the pigs are at rest
- 3 = moderate dyspnea and polypnea when the pigs
   are stressed
- 4 = moderate dyspnea and polypnea when the pigs
   are at rest
- 5 = severe dyspnea and polypnea when the pigs are stressed

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# 6 = severe dyspnea and polypnea when the pigs

In the present clinical respiratory scoring system, a score of "0" is normal, and indicates that the pig is unaffected by a porcine reproductive and respiratory disease. A score of "3" indicates moderate respiratory disease, and a score of "6" indicates very severe respiratory disease. An amount of the present vaccine or composition may be considered effective if a group of challenged pigs given the vaccine or composition show a lower average clinical respiratory score than a group of identically challenged pigs not given the vaccine or composition. (A pig is considered "challenged" when exposed to a concentration of an infectious agent sufficient to cause disease in a non-vaccinated animal.)

Preferably, the present vaccine composition is administered directly to a pig not yet exposed to a virus which causes a reproductive or respiratory disease. The present vaccine may be administered orally or parenterally. Examples of parenteral routes of administration include intradermal, intramuscular, intravenous, intraperitoneal, subcutaneous and intranasal routes of administration.

when administered as a solution, the present vaccine may be prepared in the form of an aqueous solution, a syrup, an elixir, or a tincture. Such formulations are known in the art, and are prepared by dissolution of the







antigen and other appropriate additives in the appropriate solvent systems. Such solvents include water, saline, ethanol, ethylene glycol, glycerol, Al fluid, etc. Suitable additives known in the art include certified dyes, flavors, sweeteners, and antimicrobial preservatives, such as thimerosal (sodium ethylmercurithiosalicylate). Such solutions may be stabilized, for example, by addition of partially hydrolyzed gelatin, sorbitol, or cell culture medium, and may be buffered by methods known in the art, using reagents known in the art, such as sodium hydrogen phosphate, sodium dihydrogen phosphate, potassium hydrogen phosphate and/or potassium dihydrogen phosphate.

Liquid formulations may also include suspensions and emulsions. The preparation of suspensions, for example using a colloid mill, and emulsions, for example using a homogenizer, is known in the art.

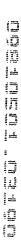
parenteral dosage forms, designed for injection into body fluid systems, require proper isotonicity and pH buffering to the corresponding levels of porcine body fluids. Parenteral formulations must also be sterilized prior to use.

Isotonicity can be adjusted with sodium chloride and other salts as needed. Other solvents, such as ethanol or propylene glycol, can be used to increase solubility of ingredients of the composition and stability of the solution. Further additives which can be used in the

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present formulation include dextrose, conventional antioxidants and conventional chelating agents, such as ethylenediamine tetraacetic acid (EDTA).

The present invention also concerns a method of producing the present vaccine, comprising the steps of synthesizing or isolating a polynucleic acid of a PRRS virus (preferably the Iowa strain) encoding an antigenic protein or portion thereof (preferably the viral coat protein), infecting a suitable host cell with the polynucleic acid, culturing the host cell, and isolating the antigenic protein or portion thereof from the culture. Alternatively, the polynucleic acid itself can confer immunoprotective activity to a host animal to which it is administered.

Preferably, the vaccine is collected from a culture medium by the steps of (i) precipitating transfected, cultured host cells, (ii) lysing the precipitated cells, and (iii) isolating the vaccine. Particularly preferably, the host cells infected with the virus or infectious agent are cultured in a suitable medium prior to collecting.

Preferably, after culturing infected host cells, the infected host cells are precipitated by adding a solution of a conventional poly(ethylene glycol) (PEG) to the culture medium, in an amount sufficient to precipitate the infected cells. The precipitated infected cells may be further purified by centrifugation. The precipitated cells

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are then lysed by methods known to those of ordinary skill in the art. Preferably, the cells are lysed by repeated freezing and thawing (three cycles of freezing and thawing is particularly preferred). Lysing the precipitated cells releases the virus, which may then be collected, preferably by centrifugation. The virus may be isolated and purified by centrifuging in a CsCl gradient, then recovering the appropriate virus-containing band from the CsCl gradient.

Alternatively, the infected cell culture may be frozen and thawed to lyse the cells. The frozen and thawed cell culture material may be used directly as a live vaccine. Preferably, however, the frozen and thawed cell culture material is lyophilized (for storage), then rehydrated for use as a vaccine.

The culture media may contain buffered saline, essential nutrients and suitable sources of carbon and nitrogen recognized in the art, in concentrations sufficient to permit growth of virus-infected cells. Suitable culture media include Dulbecco's minimal essential medium (DMEM), Eagle's minimal essential medium (MEM), Ham's medium, medium 199, fetal bovine serum, fetal calf serum, and other equivalent media which support the growth of virus-infected cells. The culture medium may be supplemented with fetal bovine serum (up to 10%) and/or L-glutamine (up to 2 mM), or other appropriate additives,

such as conventional growth supplements and/or antibiotics.

A preferred medium is DMEM.

Preferably, the present vaccine is prepared from a virus or infectious agent cultured in an appropriate cell line. The cell line is preferably PSP-36 or an equivalent cell line capable of being infected with the virus and cultured. An example of a cell line equivalent to PSP-36 is the cell line PSP-36-SAH, which was deposited under the terms of the Budapest Treaty at the American Type Culture Collection, 12301 Parklawn Drive, Rockville, Maryland 20852, U.S.A., on October 28, 1992, under the deposit number CRL 11171. Another equivalent cell line is MA-104, available commercially from Whittaker Bioproducts, Inc. (Walkersville, Maryland). Preliminary results indicate that the Iowa strain of PRRSV can also be cultured in porcine turbinate cells.

There also appears to be a relationship between the severity of histopathology caused by a challenge with a standard amount of a particular isolate and the titer to which the isolate can be grown in a mammalian host cell (e.g., CRL 11171, MA-104 cells [from African green monkey kidney], etc.).

Accordingly, the present invention also concerns a method of culturing a PRRS virus, comprising infecting cell line PSP-36, CRL 11171 or an equivalent cell line and culturing the infected cell line in a suitable medium. An

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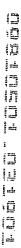
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"equivalent cell line" to PSP-36 or CRL 11171 is one which is capable of being infected with the virus and cultured, thereby producing culturable infected cells. Equivalent cell lines include MA-104, PSP-36-SAH and MARC-145 cells (available from the National Veterinary Services Laboratory, Ames, Iowa), for example.

preferably, the virus cultured is at least one isolate of the Iowa strain of PRRSV. Particularly preferably, the present vaccine is prepared from such a culture of the Iowa strain of PRRSV, cultivated in PSP-36 cells, and plaquepurified at least three times.

The cell line MA-104 is obtained from monkey kidney cells, and is epithelial-like. MA-104 cells form a confluent monolayer in culture flasks containing Dulbecco's minimal essential medium and 10% FBS (fetal bovine serum). When the monolayer is formed, the cells are inoculated with a sample of 10% homogenized tissue, taken from an appropriate tissue (such as lung and/or heart) in an infected pig. Preferably, appropriate antibiotics are present, to permit growth of virus and host cells and to suppress growth and/or viability of cells other than the host cells (e.g., bacteria or yeast).

Both PSP-36 and MA-104 cells grow some isolates of the PRRS virus to high titers (over  $10^7~TCID_{50}/ml$ ). PSP-36 and MA-104 cells will also grow the infectious agent associated



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with the Iowa strain of PRRSV. MA-104 cells also are able to grow rotaviruses, polioviruses, and other viruses.

CL2621 cells are believed to be of non-porcine origin and are epithelial-like, and are proprietary (Boehringer-Ingelheim). By contrast to PSP-36 and MA-104, some samples of the virus which causes PRRS have been unsuccessfully caltured in CL2621 cells (Bautista et al, American Association of Swine Practitioners Newsletter, 4:32, 1992).

The primary characteristics of CL2621 are that it is of non-swine origin, and is epithelial-like, growing in MEM medium. However, <u>Benfield et al</u> (*J. Vet. Diagn. Invest.*, 1992; 4:127-133) have reported that CL2621 cells were used to propagate PRRS virus, but MA-104 cells were used to control polio virus propagation, thus inferring that CL2621 is not the same as MA-104, and that the same cell may not propagate both viruses.

The Iowa strain of PRRSV generally cannot grow in cell lines other than PSP-36, PSP-36-SAH and MA-104. As described above, however, some viruses which cause PRRS have been reported to grow in both CL2621 and primary swine alveolar macrophages, although some strains of PRRS virus do not grow in PSP-36, MA-104 or CL2621 cells.

The present vaccine, virus isolates, proteins and polynucleic acids can be used to prepare antibodies which may provide immunological resistance to a patient (in this case, a pig) exposed to a virus or infectious agent.



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thereof.

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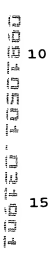




Antibodies encompassed by the present invention immunologically bind either to (1) a vaccine which protects a pig against a PRRS virus or (2) to the PRRS virus itself. The present antibodies also have the following utilities: (1) as a diagnostic agent for determining whether a pig has been exposed to a PRRS virus or infectious agent, and (2) in the preparation of the present vaccine. The present antibody may be used to prepare an immunoaffinity column by known methods, and the immunoaffinity column can be used to

isolate the virus or infectious agent, or a protein

To raise antibodies to such vaccines or viruses, one immunizes an appropriate host animal, such as a mouse, rabbit, or other animals used for such inoculation, with the protein used to prepare the vaccine. The host animal is then immunized (injected) with one of the types of vaccines described above, optionally administering an immune-enhancing agent (adjuvant), such as those described The host animal is preferably subsequently immunized from 1 to 5 times at certain intervals of time, preferably every 1 to 4 weeks, most preferably every 2 The host animals are then sacrificed, and their blood is collected. Sera is then separated by known techniques from the whole blood collected. The sera contains antibodies to the vaccines. Antibodies can also



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be purified by known methods to provide immunoglobulin G (IqG) antibodies.

The present invention also encompasses monoclonal antibodies to the present vaccines and/or viruses. Monoclonal antibodies may be produced by the method of Kohler et al (Nature, vol. 256 (1975), pages 495-497). Basically, the immune cells from a whole cell preparation of the spleen of the immunized host animal (described above) are fused with myeloma cells by a conventional procedure to produce hybridomas. Hybridomas are cultured, and the resulting culture fluid is screened against the fluid or inoculum carrying the infectious agent (virus or Introducing the hybridoma into the peritoneum of vaccine). the host animal produces a peritoneal growth of the Collection of the ascites fluid of the host hybridoma. animal provides a sample of the monoclonal antibody to the infectious agent produced by the hybridoma. supernatant from the hybridoma cell culture can be used as a source of the monoclonal antibody, which is isolated by methods known to those of ordinary skill in the art. Preferably, the present antibody is of the IgG or IgM type of immunoglobulin.

The present invention also concerns a method of treating a pig suffering from a reproductive and respiratory disease, comprising administering an effective amount of an antibody which immunologically binds to a



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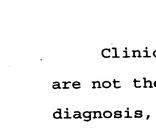




virus which causes a porcine reproductive and respiratory disease or to a vaccine which protects a pig against infection by a porcine reproductive and respiratory virus in a physiologically acceptable carrier to a pig in need thereof.

The present method also concerns a method of diagnosing infection of a pig by or exposure of a herd to a porcine reproductive and respiratory syndrome virus and a diagnostic kit for assaying the same, comprising the present antibody (preferably a monoclonal antibody) and a diagnostic agent which indicates a positive immunological reaction with said antibody (preferably comprising peroxidase-conjugated streptavidin, a biotinylated antibody to a PRRSV protein or antigen and a peroxidase). The present kit may further comprise aqueous hydrogen peroxide, a protease which digests the porcine tissue sample, a fluorescent dye (e.g., 3,3'-diaminobenzidine tetrahydrochloride), and a tissue stain (e.g., hematoxylin).

A diagnosis of PRRS relies on compiling information from the clinical history of the herd being diagnosed, from serology and pathology of infected pigs, and ultimately, on isolation of the PRRS virus (PRRSV) from the infected herd. Thus, the present method of detecting PRRSV is useful in diagnosing infection by and/or exposure to the virus in a herd.



Clinical signs vary widely between farms, and thus, are not the most reliable evidence of a definitive diagnosis, except in the case of a severe acute outbreak in naive herds which experience abortion storms, increased numbers of stillborn pigs, and severe neonatal and nursery pig pneumonia. Presently, the most common clinical presentation is pneumonia and miscellaneous bacterial problems in 3-10 week old pigs. However, many PRRSV-positive herds have no apparent reproductive or respiratory problems.

There are some gross lesions that are very suggestive of PRRSV infection in growing pigs. The most consistent experimentally reproducible gross lesion in 3-10 week-old pigs inoculated with several different PRRSV strains is lymphadenopathy. In particular, iliac and mediastinal lymph nodes are often 3-10 times normal size, tan in color, and sometimes cystic. The lymph nodes are not normally hyperemic, such as the lesion/conditions seen in bacterial septicemia.

Three histologic lesions are consistent with PRRSV infection. Interstitial pneumonia is commonly observed and is characterized by septal infiltration with mononuclear cells, type 2 pneumocyte proliferation, and the presence of necrotic cells in the alveolar spaces. Nonsuppurative perivascular myocarditis and hyperplastic lymph nodes are commonly observed in the subacute stages of disease.

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The degree of grossly visible pneumonia is strain dependent. In general, the lungs fail to collapse and have a patchy distribution of 10-80% tan-colored consolidation with irregular borders. Encephalitis is less often observed. Lesions in the fetus and placenta are rarely observed by light microscopy.

However, the percentage of consolidation in the lungs provides a particularly reliable test for infection by PRRSV (i.e.,  $\geq$  10% consolidation at any time from 3 to 10 days post-infection (DPI) is a positive indication of infection), particularly by a high virulence phenotype (hv) virus ( $\geq$  40% consolidation at any time from 3 to 10 days DPI is a positive indication of infection by an hv PRRSV isolate).

In contrast to histopathology on lung tissue(s), most laboratories are routinely using either an indirect-fluorescent antibody (IFA) test or immunoperoxidase monolayer assay (IPMA) for serum antibody detection. With both the IFA and IPMA, one must subjectively determine endpoints and thus the tests are not automatable. Serum virus (SVN) neutralization tests have also been developed, and ELISA tests are currently used in some research laboratories. Antibodies detected by the IFA test usually appear with 10 days of exposure but may be relatively short-lived, sometimes disappearing within 3 months.

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Antibodies detected by ELISA usually appear within 3 weeks, but their duration is unknown. SVN antibodies usually are not detected until 4-5 weeks post exposure. The SVN test is considered less sensitive in acute disease, but improvements have been made in the SVN test using seronegative porcine serum supplementation. SVN titers reportedly are measurable longer than titers in IFA and IPMA, and thus, may be better suited for detection of positive animals in chronically infected herds.

In IFA, infected cells are fixed with acetone and methanol solutions, and antibodies for the convalescent sera of infected pigs are incubated with the infected cells, preferably for about 30 min. at 37°C. A positive immunological reaction is one in which the antibody binds to the virus-infected cells, but is not washed out by subsequent washing steps (usually 3 X with PBS buffer). A second antibody (an anti-antibody) labeled with a fluorescent reagent (FITC) is then added and incubated, preferably for anther 30 min. A positive immunological reaction results in the second antibody binding to the first, being retained after washing, and resulting in a fluorescent signal, which can be detected and semiquantified. A negative immunological reaction results in little or no binding of the antibody to the infected cell. Therefore, the second, fluorescently-labeled antibody fails to bind, the fluorescent label is washed out, and little or

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no fluorescence is detected, compared to an appropriate positive control.

IPA and ELISA kits are similar to the IFA kit, except that the second antibody is labeled with a specific enzyme, instead of a fluorescent reagent. Thus, one adds an appropriate substrate for the enzyme bound to the second antibody which results in the production of a colored product, which is then detected and quantified by colorimetry, for example.

Clinicians use antibody titers to determine the appropriate time for vaccination and/or implementation of management or control strategies. Prior to the present invention, serology tests did not provide antibody titer levels adequate or reliable enough to make animal health care decisions. It may have been appropriate to look for a change from seronegative to seropositive status, or for at least a 4-fold increase in titer, as a positive indication of PRRSV infection/exposure. Looking for an increasing percentage of seropositive pigs in a particular age group over time in a herd can be useful to determine where the virus is maintained and actively spreading. Sows infected in the early 3rd trimester and aborting near term will likely not show increasing titers, however.

Virus isolation (VI) provides a definitive diagnosis, but has some limitations. Virus is rarely isolated from stillborn or autolyzed aborted fetuses. Sows infected

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early in the last trimester may have transient viremia and not abort until late term. Dead pigs of any age are not the best samples for VI, because the virus does not survive well at room temperature. Tissues should be removed from the carcass, packaged separately, and refrigerated as soon as possible to obtain a viable virus sample.

The best tissues for virus isolation are tonsil, lung, lymph nodes, and spleen. Serum is also an excellent sample for virus isolation, since (a) viremia is often prolonged in growing pigs, (b) the sample is easy to handle, and (c) the sample can be quickly chilled and processed.

Variation between laboratories in the ability to isolate PRRSV is high because the tests, reagents, cell lines, and media used to detect/screen for PRRSV have not been standardized. The efficacy of isolation varies because not all North American strains will grow on each cell line. Frozen tissue-section IFA tests have been used with limited success.

Serum virus neutralization (SVN) tests have also been developed, and ELISA tests are currently used in some research laboratories. Antibodies detected by ELISA usually appear within 3 weeks, but their duration is unknown. SVN antibodies usually are not detected until 4-5 weeks post-exposure. The SVN test is considered less sensitive in acute disease, but improvements have been made in the SVN test using seronegative porcine serum

supplementation. SVN titers reportedly are measurable for a longer period of time than titers in IFA and IPMA. Thus, SVN titers may be better suited for detection of positive animals in chronically infected herds.

Prior to the present invention, however, serology tests did not provide antibody titer levels adequate or reliable enough to make animal health care decisions. Looking for an increasing percentage of seropositive pigs in a particular age group over time in a herd can also be useful to determine where the virus is maintained and actively spreading. Sows infected in the early third trimester and aborting near term will likely not show increasing titers, however. Thus, although it may have been appropriate to look for a change from seronegative to seropositive status or for at least a 4-fold increase in titer as a positive indication of PRRSV infection and/or exposure, a need for a more reliable titer-based assay is felt.

Thus, the present invention also concerns a method for detecting PRRSV antigen in tissues. The present diagnostic method, employing an immunoperoxidase test (IPT) preferably on formalin-fixed tissue, appears to be quite useful to confirm the presence of active infection, and may provide a significant and meaningful increase in the reliability of titer-based assays. A section of lungs, tonsils, mediastinal lymph nodes, and tracheobronchial lymph nodes

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from 26 pigs experimentally inoculated with ATCC VR 2385 PRRSV was examined (see Experiment V below). The virus was detected in 18/26 lungs, 26/26 tonsils, 15/26 mediastinal lymph nodes, and 14/26 tracheobronchial lymph nodes. pigs in this study were killed over a 28 day period (postinoculation). The virus was detected in at least one tissue in every pig necropsied up to 10 days post inoculation.

A complete technique for the present immunoperoxidase technique for PRRSV antigen detection in porcine tissues, based on a streptavidin-biotin assay, is described in Example V hereinunder. Briefly, the present method for detecting PRRSV comprises removing endogenous peroxidase from an isolated porcine tissue sample with aqueous hydrogen peroxide (preferably, a 0.1-5%, and more preferably, 0.1-1.0% solution), then digesting the tissue with sufficient amount of an appropriate protease to expose viral antigens (for example, Protease XIV, Sigma Chemical Company, St. Louis, MO, and more preferably, a 0.001-0.25% Thereafter, the method further aqueous solution thereof). comprises incubating primary monoclonal antibody ascites fluid (preferably diluted in TRIS/PBS by an amount of from 1:10 to 1:100,000, and more preferably, from 1:100 to 1:10,000) with the protease-treated tissue sections in a humidified chamber for a sufficient length of time and at an appropriate temperature to provide essentially complete

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immunological binding to occur, if it can in fact occur (e.g., 16 hours at 4°C).

One suitable monoclonal antibody for use in the present diagnostic assay is SDOW-17 (available from Dr. David Benfield, South Dakota State Univ.), which recognizes a conserved epitope of the PRRSV nucleocapsid protein (Nelson et al, "Differentiation of U.S. and European isolates of porcine reproductive and respiratory syndrome virus by monoclonal antibodies," J. Clin. Micro., 31:3184-3189 (1993)).

The present method for detecting PRRSV then further comprises incubating biotinylated goat anti-mouse linking antibody (available from Dako Corporation, Carpintera, CA) with the tissue, followed by incubating peroxidase-conjugated streptavidin with the biotinylated antibody-treated tissue (Zymed Laboratories, South San Francisco, CA). The method then further comprises incubating the peroxidase-conjugated streptavidin-treated tissue with a chromagen, such as 3,3'-diaminobenzidine tetrahydrochloride (available from Vector Laboratories Inc., Burlingame, CA), and finally, staining the treated tissue with hematoxylin.

particularly when combined with the further diagnostic techniques of histopathology, virus isolation procedures and serology, the present tissue immunoperoxidase antigen detection technique offers a rapid and reliable diagnosis of PRRSV infection.

Other features of the invention will become apparent in the course of the following descriptions of exemplary embodiments, which are given for illustration of the invention, and are not intended to be limiting thereof.

EXPERIMENT I

MOLECULAR CLONING AND NUCLEOTIDE SEQUENCING OF THE 3'-TERMINAL REGION OF VR 2385 (PLAQUE-PURIFIED ISU-12)

(I) Materials and Methods

(A) Virus Propagation and Purification

A continuous cell line, PSP-36, was used to isolate and propagate ISU-12. The ISU-12 virus was plaque-purified 3 times on PSP-36 cells (plaque-purified ISU-12 virus was deposited under the terms and conditions of the Budapest Treaty at the American Type Culture Collection, 12301 Parklawn Drive, Rockville, Maryland 20852, U.S.A., under Accession No. VR 2385). The PSP-36 cells were then infected with the plaque-purified virus. When more than 70% of the infected cells showed cytopathic changes, the culture was frozen and thawed three times. medium was then clarified by low-speed centrifugation at The virus was then 5,000 X g for 15 min. at 4°C. precipitated with 7% PEG-8000 and 2.3% NaCl at 4°C overnight with stirring, and the precipitate was pelleted by centrifugation. The virus pellets were then resuspended in 2 ml of tris-EDTA buffer, and layered on top of a CsCl

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gradient (1.1245-1.2858 g/ml). After ultracentrifugation at 28,000 rpm for about 8 hours at 20°C, a clear band with a density of 1.15-1.18 g/ml was observed and harvested. The infectivity titer of this band was determined by IFA, and the titer was found to be 10<sup>6</sup> TCID<sub>50</sub>/ml. Typical virus particles were also observed by negative staining electron microscopy (EM).

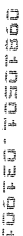
## (B) Isolation of Viral RNA

Total RNA was isolated from the virus-containing band in the CsCl gradient with a commercially available RNA isolation kit (obtained from Stratagene). Poly(A) RNA was then enriched by oligo (dT)-cellulose column chromatography according to the procedure described by the manufacturer of the column (Invitrogen).

# (C) Construction of VR 2385 cDNA λ library

A general schematic procedure for the construction of a cDNA  $\lambda$  library is shown in Figure 3. First strand cDNA synthesis from mRNA was conducted by reverse transcription using an oligo (dT) primer having a Xho I restriction site. The nucleotide mixture contained normal dATP, dGTP, dTTP and the analog 5-methyl dCTP, which protects the cDNA from restriction enzymes used in subsequent cloning steps.

Second strand cDNA synthesis was then conducted with RNase H and DNA polymerase I. The cDNA termini were



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blunted (blunt-ended) with T4 DNA polymerase, ligated to ECOR I adaptors with T4 DNA ligase, and subsequently phosphorylated with T4 polynucleotide kinase. The cDNA was digested with Xho I, and the digested cDNA were size-selected on an agarose gel. Digested cDNA larger than 1 kb in size were selected and purified by a commercially available DNA purification kit (GENECLEAN, available from BIO 101, Inc., La Jolla, California).

The purified cDNA was then ligated into lambda phage vector arms, engineered with Xho I and EcoR I cohesive ends. The ligated vector was packaged into infectious lambda phages with lambda extracts. The SURE strain (available from Stratagene) of *E. coli* cells were used for transfection, and the lambda library was then amplified and titrated in the XL-1 blue cell strain.

(D) Screening the  $\lambda$  Library by Differential Hybridization

A general schematic procedure for identifying authentic clones of the PRRS virus VR 2385 strain by differential hybridization is shown in Figure 4, and is described hereunder. The  $\lambda$  library was plated on XL-1 blue cells, plaques were lifted onto nylon membranes in duplicates, and denatured with 0.5 N NaOH by conventional methodology. Messenger RNA's from both virus-infected PSP-36 cells and non-infected PSP-36 cells were isolated by

oligo (dT) cellulose column chromatography as described by the manufacturer of the column (Invitrogen).

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Complementary DNA probes were synthesized from mRNA's isolated from virus-infected PSP-36 cells and normal PSP-36 cells using random primers in the presence of 32P-dCTP according to the procedure described by the manufacturer (Amersham). Two probes (the first synthesized from virusinfected PSP-36 cells, the other from normal, uninfected PSP-36 cells) were then purified individually by Sephadex G-50 column chromatography. The probes were hybridized with the duplicated nylon membranes, respectively, at 42°C Plaques which hybridized with the probe in 50% formamide. prepared from virus infected cells, but not with the probe prepared from normal cells, were isolated. The phagemids containing viral cDNA inserts were rescued by in vitro excision with the help of G408 helper phage. The rescued phagemids were then amplified on XL-1 blue cells. plasmids containing viral cDNA inserts were isolated by Qiagen column chromatography, and were subsequently sequenced.

(E) Nucleotide Sequencing and Sequence Analysis

Plasmids containing viral cDNA inserts were purified
by Qiagen column chromatography, and sequenced by Sanger's
dideoxy method with universal and reverse primers, as well
as a variety of internal oligonucleotide primers.

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Sequences were obtained from at least three separate clones. Additional clones or regions were sequenced when ambiguous sequence data were obtained. The nucleotide sequence data were assembled and analyzed independently using two computer software programs, GENEWORKS (IntelliGenetics, Inc., Mountain View, California) and MACVECTOR (International Biotechnologies, Inc., New Haven, Connecticut).

### (F) Oligonucleotide Primers

Oligonucleotides were synthesized as single-stranded DNA using an automated DNA synthesizer (Applied Biosystems) and purified by HPLC. Oligonucleotides PP284 (5'-CGGCCGTGTG GTTCTCGCCA AT-3'; SEQ ID NO:1) and PP285 (5'-CCCCATTTCC CTCTAGCGAC TG-3'; SEQ ID NO:2) were synthesized for PCR amplification. A DNA probe was generated with these two primers from the extreme 3' end of the viral genome for Northern blot analysis (see discussion below). Oligonucleotides PP286 (5'-GCCGCGGAAC CATCAAGCAC-3'; SEQ ID NO:3) and PP287 (5'-CAACTTGACG CTATGTGAGC-3'; SEQ ID NO:4)were synthesized for PCR amplification. A DNA probe generated by these two primers was used to further screen the  $\lambda$  library. Oligonucleotides PP288 (5'-GCGGTCTGGA TTGACGACAG-3'; SEQ ID NO:5), PP289 (5'-GACTGCTAGG GCTTCTGCAC-3'; SEQ ID NO:6), PP386 (5'-GCCATTCAGC

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TCACATAGCG-3'; SEQ ID NO:7), PP286 and PP287 were used as sequencing primers to obtain internal sequences.

## Northern Blot Analysis

A specific DNA fragment from the extreme 3' end of the VR 2385 cDNA clone was amplified by PCR with primers PP284 The DNA fragment was excised from an agarose and PP285. gel with a commercially available DNA purification kit (GENECLEAN, obtained from Bio 101), and labeled with 32P-dCTP by random primer extension (using a kit available from Amersham). Total RNA was isolated from VR 2385infected PSP-36 cells at 36 hours post-infection, using a commercially available kit for isolation of total RNA according to the procedure described by the manufacturer (Stratagene). VR 2385 subgenomic mRNA species were denatured with 6 M glyoxal and DMSO, and separated on a 1% agarose gel. (Results from a similar procedure substituting a 1.5% agarose gel are described in Experiment II below and are shown in Figure 5.) The separated subgenomic mRNA's were then transferred onto nylon membranes using a POSIBLOT pressure blotter (Stratagene). Hybridization was carried out in a hybridization oven with roller bottles at 42°C and 50% formamide.

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# RESULTS

(A) Cloning, Identification and Sequencing of VR 2385 3' Terminal Genome

An oligo (dT)-primed cDNA λ library was constructed from a partially purified virus, obtained from VR 2385infected PSP-36 cells. Problems were encountered in screening the cDNA  $\lambda$  library with probes based on the Lelystad virus sequence. Three sets of primers were The first set (PP105 and PP106; SEQ ID NOS:8-9) correspond to positions 14577 to 14596 and 14977 to 14995 of the Lelystad genomic sequence, located in the The second set (PP106 and PP107, nucleocapsid gene region. SEQ ID NOS:9-10) correspond to positions 14977 to 14995 and 14054 to 14072 of the Lelystad genomic sequence, flanking ORF's 6 and 7. The third set (PM541 and PM542; SEQ ID NOS:11-12) correspond to positions 11718 to 11737 and 11394 to 11413 of the Lelystad genomic sequence, located in the ORF-1b region.

PP105: 5'-CTCGTCAAGT ATGGCCGGT-3' (SEQ ID NO:8)

PP106: 5'-GCCATTCGCC TGACTGTCA-3' (SEQ ID NO:9)

PP107: 5'-TTGACGAGGA CTTCGGCTG-3' (SEQ ID NO:10)

PM541: 5'-GCTCTACCTG CAATTCTGTG-3' (SEQ ID NO:11)

PM542: 5'-GTGTATAGGA CCGGCAACCG-3' (SEQ ID NO:12)

All attempts to generate probes by PCR from the VR
25 2385 infectious agent using these three sets of primers

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were unsuccessful. After several attempts using the differential hybridization technique, however, the authentic plaques representing VR 2385-specific cDNA were isolated using probes prepared from VR 2385-infected PSP-36 cells and normal PSP-36 cells. The procedures involved in differential hybridization are described and set forth in Figure 4.

Three positive plaques ( $\lambda$ -4,  $\lambda$ -75 and  $\lambda$ -91) were initially identified. Phagemids containing viral cDNA inserts within the  $\lambda$  phage were rescued by in vitro excision with the help of G408 helper phages. The inserts of the positive clones were analyzed by restriction enzyme digestion and terminal sequencing. The specificity of the cDNA clones was further confirmed by hybridization with RNA from PSP-36 cells infected with the Iowa strain of PRRSV, but not with RNA from normal PSP-36 cells. A DNA probe was then generated from the 5'-end of clone  $\lambda$ -75 by PCR with primers PP286 and PP287. Further positive plaques ( $\lambda$ -229,  $\lambda$ -268,  $\lambda$ -275,  $\lambda$ -281,  $\lambda$ -323 and  $\lambda$ -345) were identified using this probe. All  $\lambda$  cDNA clones used to obtain the 3'terminal nucleotide sequences are presented in Fig. 6. least three separate clones were sequenced to eliminate any In the case of any ambiguous sequence data, mistakes. additional clones and internal primers (PP288, PP289, PP286, PP287 and PP386) were used to determine the The 2062-bp 3'-terminal sequence (SEQ ID NO:13) sequence.

and the amino acid sequences/encoded by ORF's 5, 6 and 7 (SEQ ID NOS:15, 17 and 19, respectively) are presented in Figure 7.

# (B) A Nested Set of Subgenomic mRNA

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Total RNA from virus-infected PSP-36 cells was separated on 1% glyoxal/DMSO agarose gel, and blotted onto nylon membranes. A cDNA probe was generated by PCR with a set of primers (PP284 and PP285) flanking the extreme 3'-terminal region of the viral genome. The probe contains a 3'-nontranslational sequence and most of the ORF-7 sequence. Northern blot hybridization results show that the pattern of mRNA species from PSP-36 cells infected with the Iowa strain of PRRSV is very similar to that of Lelystad virus (LV), equine arteritis virus (EAV), lactate dehydrogenase-elevating virus (LDV) and coronavirus, in that virus replication required the formation of subgenomic mRNA's.

The results also indicate that VR 2385-specific subgenomic mRNA's represent a 3'-nested set of mRNA's, because the Northern blot probe represents only the extreme 3' terminal sequence. The size of VR 2385 viral genomic RNA (14 kb) and 6 subgenomic mRNA's (RNA 2 (3.0 kb), RNA 3 (2.5 kb), RNA 4 (2.2 kb), RNA 5 (1.8 kb), RNA 6 (1.3 kb) and RNA 7 (0.98 kb)) resemble those of LV, although there are differences in both the genome and in subgenomic RNA

(C) Analysis of Open Reading Frames Encoded by Subgenomic RNA

Three large ORF's have been found in SEQ ID NO:13: ORF-5 (nucleotides [nt] 426-1025; SEQ ID NO:14), ORF 6 (nt 1013-1534; SEQ ID NO:16) and ORF 7 (nt 1527-1895; SEQ ID NO:18). ORF 4, located at the 5' end of the resulting sequence, is incomplete in the 2062-bp 3'-terminal sequence ORF'S 5, 6 AND 7 each have a coding of SEQ ID NO:13. capacity of more than 100 amino acids. ORF 5 and ORF 6 overlap each other by 13 bp, and ORF 6 and ORF 7 overlap each other by 8 bp. Two smaller ORF's located entirely within ORF 7 have also been found, coding for only 37 aa and 43 aa, respectively. Another two short ORF's overlap fully with ORF 5. The coding capacity of these two ORF's is only 29 aa and 44 aa, respectively. No specific subgenomic mRNA's were correlated to these smaller ORF's by Northern blot analysis. ORF 6 and ORF 7 are believed to encode the viral membrane protein and capsid protein, respectively.

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- (D) Consensus Sequence for Leader Junction
  Sequence analysis shows that a short sequence motif,

  AACC, may serve as the site in the subgenomic mRNA's where
  the leader is added during transcription (the junction
  site). The junction site of ORF 6 is found 21 bp upstream
  from the ATG start codon, and the junction site of ORF 7 is
  found 13 bp upstream from the ATG start codon,
  respectively. No AACC consensus sequence has been
  identified in ORF 5, although it has been found in ORF 5 of
  LV. Similar junction sequences have been found in LDV and
  EAV.
  - (E) 3'-Nontranslational Sequence and Poly (A) Tail
    A 151 nucleotide-long (151 nt) nontranslational
    sequence following the stop codon of ORF 7 has been
    identified in the genome of VR 2385, compared to 114 nt in
    LV, 80 nt in LDV and 59 nt in EAV. The length of the poly
    (A) tail is at least 13 nucleotides. There is a consensus
    sequence, CCGG/AAATT-poly (A) among PRRS virus VR 2385, LV
    and LDV in the region adjacent to the poly (A) tail.
    - (F) Sequence Comparison of VR 2385 and LV Genomes
      Among ORF's 5, 6 and 7, and Among the
      Nontranslational Sequences

A comparison of the ORF-5 regions of the genomes of VR 2385 and of the Lelystad virus (SEQ ID NO:20) is shown in

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Figure 8. The corresponding comparisons of the ORF-6 region, the ORF-7 region, and the nontranslational sequences of VR 2385 (SEQ ID NOS:16, 18 and 22, respectively) with the corresponding regions of LV (SEQ ID NOS:23, 25 and 27, respectively) are shown in Figures 9, 10 and 11, respectively.

The results of the comparisons are presented in Table 1 below. The nucleotide sequence homologies between LV and VR 2385 of the ORF 5, ORF 6, ORF 7 and the nontranslational sequences are 53%, 78%, 58% and 58%, respectively.

The size of ORF 7 in LV is 15 nt larger than that in VR 2385. Also, the 3'-terminal nontranslational sequence is different in length (150 nt in VR 2385, but only 114 nt in LV). Like LV, the junction sequence, AACC, has also been identified in the genome of the Iowa strain of PRRS virus isolate VR 2385, except for ORF 5. The junction sequence of ORF 6 in VR 2385 is 21 nt upstream from the ATG start codon, whereas the junction sequence of ORF 6 is 28 nt upstream from ATG in LV.





Table 1: Comparison of genes of U.S. PRRSV isolate ATCC VR 2385 with those of European isolate Lelystad virus\*

Gene	RNA	Estimated RNA size	ORFs	VR 2385			Lelystad			Homology between
		(in Kb)		Size amino acids	N-glyco- sylation sites	Pred. protein size (kd)	Size amino acids	N-glyco- sylation sites	Pred. protein size (kd)	VR 2385 & Lelystac
5	5	1.9	5	200	2	22.2	201	2	22.4	53
6	6	1.4	6 -	174	1	19.1	173	2	18.9	78
7	7	0.9	7	123	. 2	13.6	128	1	13.8	58
NTR	•	-	-	151 (nt)	<b>-</b>	NA NA	114 (nt)	0	NA	58 (nt)

\*: Based on data presented by <u>Conzelmann et al</u>, *Virology*, 193, 329-339 (1993), <u>Meulenberg et al</u>, *Virology*, 192, 62-72 (1993), and the results presented herein.

#### EXPERIMENT II

THE EXPRESSION OF VR 2385 GENES IN INSECT CELLS

(A) Production of Recombinant Baculovirus

The ORF-5, ORF-6 and ORF-7 sequences were individually amplified by PCR using primers based on the VR 2385

(ISU-12) genomic nucleotide sequence. ORF-5 was amplified using the following primers:

- 5'-GGGGATCCGG TATTTGGCAA TGTGTC-3' (SEQ ID NO:28)
- 3'-GGGAATTCGC CAAGAGCACC TTTTGTGG-5' (SEQ ID NO:29)

ORF-6 was amplified using the following primers:

5'-GGGGATCCAG AGTTTCAGCG G-3' (SEQ ID NO:30)

[=<u>}</u>





3'-GGGAATTCTG GCACAGCTGA TTGAC-5' (SEQ ID NO:31)
ORF-7 was amplified using the following primers:

5'-GGGGATCCTT GTTAAATATG CC-3' (SEQ ID NO:32)
3'-GGGAATTCAC CACGCATTC-5' (SEQ ID NO:33)

The amplified DNA fragments were cloned into baculovirus transfer vector pVL1393 (available from Invitrogen). One  $\mu g$  of linearized baculovirus AcMNPV DNA (commercially available from Pharmingen, San Diego, California) and 2  $\mu g$  of PCR-amplified cloned cDNA-containing vector constructs were mixed with 50  $\mu l$  of lipofectin (Gibco), and incubated at 22°C for 15 min. to prepare a transfection mixture.

One hour after seeding HI-FIVE cells, the medium was replaced with fresh Excell 400 insect cell culture medium (available from JR Scientific Co.), and the transfection mixture was added drop by drop. The resulting mixture was incubated at 28°C for six hours. Afterwards, the transfection medium was removed, and fresh Excell 400 insect cell culture medium was added. The resulting mixture was then incubated at 28°C.

Five days after transfection, the culture medium was collected and clarified. Ten-fold dilutions of supernatants were inoculated onto HI-FIVE cells, and incubated for 60 min. at room temperature. After the inoculum was discarded, an overlay of 1.25% of agarose was

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applied onto the cells. Incubation at 28°C was conducted for four days. Thereafter, clear plaques were selected and picked using a sterile Pasteur pipette. Each plaque was mixed with 1 ml of Grace's insect medium into a 5 ml snap cap tube, and placed in a refrigerator overnight to release the virus from the agarose. Tubes were centrifuged for 30 minutes at 2000 x g to remove agarose, and the supernatants were transferred into new sterile tubes. Plaque purification steps were repeated three times to avoid possible wild-type virus contamination. Pure recombinant clones were stored at -80°C for further investigation.

(B) Expression of Recombinant Iowa Strain Infectious
Agent Proteins

Indirect immunofluorescence assay and radioimmunoprecipitation tests were used to evaluate expression.

Indirect immunofluorescence assay: Hi-five insect cells in a 24-well cell culture cluster plate were infected with wild-type baculovirus or recombinant baculovirus, or were mock-infected. After 72 hours, cells were fixed and stained with appropriate dilutions of swine anti-VR 2385 polyclonal antibodies, followed by fluorescein isothiocyanate-labelled (FITC-labelled) anti-swine IgG. Immunofluorescence was detected in cells infected with the recombinant viruses, but not in mock-infected cells or

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cells inoculated with wild-type baculovirus. For example, Figure 12 shows HI-FIVE cells infected with the recombinant baculovirus containing the VR 2385 ORF-7 gene (Baculo.PRRSV.7), which exhibit a cytopathic effect. Similar results were obtained with recombinant baculovirus containing ORF-5 (Baculo.PRRSV.5) and ORF-6 (Baculo.PRRSV.6; data not shown). Figures 13 and 14 show HI-FIVE cells infected with a recombinant baculovirus containing the VR 2385 ORF-6 gene and VR 2385 ORF-7 gene, respectively, stained with swine antisera to VR 2385, followed by fluorescein-conjugated anti-swine IgG, in which the insect cells are producing recombinant Iowa strain viral protein. Similar results were obtained with recombinant baculovirus containing ORF-5.

Radioimmunoprecipitation: Radioimmunoprecipitation was carried out with each recombinant virus (Baculo.PRRSV.5, Baculo.PRRSV.6 and Baculo.PRRSV.7) to further determine the antigenicity and authenticity of the recombinant proteins. HI-FIVE insect cells were mockinfected, or alternatively, infected with each of the recombinant baculoviruses. Two days after infection, methionine-free medium was added. Each mixture was incubated for two hours, and then proteins labeled with <sup>35</sup>S-methionine (Amersham) were added, and the mixture was incubated for four additional hours at 28°C. Radiolabeled cell lysates were prepared by three cycles of freezing and

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thawing, and the cell lysates were incubated with preimmune or immune anti-VR 2385 antisera. The immune complexes were precipitated with Protein A agarose and analyzed on SDS-PAGE after boiling. X-ray film was exposed to the gels at -80°C, and developed. Bands of expected size were detected with ORF-6 (Figure 15) and ORF-7 (Figure 16) products.

### EXPERIMENT III

### Summary:

The genetic variation and possible evolution of porcine reproductive and respiratory syndrome virus (PRRSV) was determined by cloning and sequencing the putative membrane protein (M, ORF 6) and nucleocapsid (N, ORF 7) genes of six U.S. PRRSV isolates with differing virulence. The deduced amino acid sequences of the putative M and N proteins from each of these isolates were aligned with the corresponding sequences (to the extent known) of one other U.S. isolate, two European isolates, and other members of the proposed arterivirus group, including lactate dehydrogenase-elevating virus (LDV) and equine arteritis virus (EAV).

The putative M and N genes displayed 96-100% amino acid sequence identity among U.S. PRRSV isolates with differing virulence. However, their amino acid sequences varied extensively from those of European PRRSV isolates,

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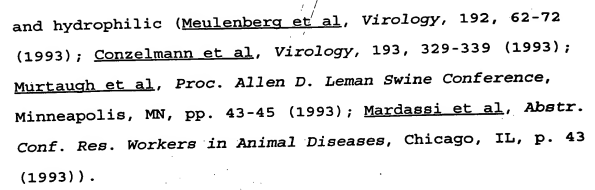
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and displayed only 57-59% and 78-81% identity, respectively. The U.S. PRRSV isolates were more closely related to LDV than were the European PRRSV isolates. The N protein of the U.S. isolates and European isolates shared about 50% and 40% amino acid sequence identity with that of LDV, respectively.

The phylogenetic dendrograms constructed on the basis of the putative M and N genes of the proposed arterivirus group were similar and indicated that both U.S. and European PRRSV isolates were related to LDV and were distantly related to EAV. The U.S. and European PRRSV isolates fell into two distinct groups with slightly different genetic distance relative to LDV. The results suggest that U.S. and European PRRSV isolates represent two different genotypes, and that they may have evolved from LDV at different time periods and have existed separately in U.S. and Europe before their association with PRRS was recognized in swine.

ORF 6 encodes the membrane protein (M) of PRRSV, based on the similar characteristics of the ORF 6 of EAV, ORF 2 of LDV, and the M protein of mouse hepatitis virus and infectious bronchitis virus (Meulenberg et al, Virology, 192, 62-72 (1993); Conzelmann et al, Virology, 193, 329-339 (1993); Mardassi et al, Abstr. Conf. Res. Workers in Animal Diseases, Chicago, IL, p. 43 (1993)). The product of ORF 7, the viral nucleocapsid protein (N), is extremely basic

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The amino acid sequences encoded by ORF's 5, 6 and 7 of U.S. isolate VR 2385 and of the European isolate

Lelystad virus (LV) have been compared, and the identity

(i.e., the percentage of amino acids in sequence which are the same) between the two viruses is only 54%, 78% and 58%, respectively. Thus, striking genetic differences exist between the U.S. isolate VR 2385 and the European isolate LV (see U.S. application Serial No. 08/131,625, filed October 5, 1993).

However, the U.S. isolate VR 2385 is highly pathogenic compared to European LV. Thus, PRRSV isolates in North America and in Europe appear to be antigenically and genetically heterogeneous, and different genotypes or serotypes of PRRSV may exist.

To further determine the genetic variation among the PRRSV isolates, the putative M and N genes of five additional U.S. PRRSV isolates with differing virulence were cloned and sequenced. Phylogenetic trees based on the putative M and N genes of seven U.S. PRRSV isolates, two European PRRSV isolates and other members of the proposed

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arterivirus group, including LDV and EAV, have been constructed.

PRRSV isolates (ISU-12 (VR 2385/VR 2386), ISU-22 (VR 2429), ISU-55 (VR 2430), ISU-79, ISU-1894 and ISU-3927 (VR 2431), each of which is disclosed and described in U.S. application Serial No. 08/131,625, filed October 5, 1993) were isolated from pig lungs obtained from different farms in Iowa during PRRS outbreaks, according to the procedure described in U.S. application Serial No. 08/131,625. A continuous cell line, ATCC CRL 11171, was used to isolate and propagate these viruses. All viruses were biologically cloned by three cycles of plaque purification prior to polynucleic acid sequencing.

Pathogenicity studies in caesarean-derived colostrum-deprived (CDCD) pigs, described in U.S. application Serial No. 08/131,625, showed that VR 2385, VR 2429 and ISU-79 were highly pathogenic, whereas VR 2430, ISU-1894 and VR 2431 were not as pathogenic. For example, VR 2385, VR 2429 and ISU-79 produced from 50 to 80% consolidation of the lung tissues in experimentally-infected five-week-old CDCD pigs necropsied at 10 days post inoculation, whereas VR 2430, ISU-1894 and VR 2431 produced only 10 to 25% consolidation of lung tissues in the same experiment.





### Experimental Section:

Monolayers of ATCC CRL 11171 cells were infected with each of the PRRSV isolates at the seventh passage at an m.o.i. of 0.1. Total cellular RNA was isolated from infected cells by the guanidine isothiocyanate method (Sambrook et al, "Molecular Cloning: A Laboratory Manual," 2nd ed., Cold Spring Harbor Laboratory, Cold Spring Harbor, New York (1989)). The quality of RNA from each isolate was determined by Northern blot hybridization (data not shown) with a cDNA probe generated from the extreme 3'-end of the VR 2385 genome by the polymerase chain reaction (PCR) with primers PP284 and PP285 (SEQ ID NOS: 1 AND 2), as described in U.S. Application Serial No. 08/131,625. cDNA was synthesized from total cellular RNA with random primers using reverse transcriptase. The synthesized cDNA was amplified by polymerase chain reaction (PCR) as described previously (Meng et al, J. Vet. Diagn. Invest., 5, 254-258 (1993)). Primers for RT-PCR were designed on the basis of a sequence in the genome of VR 2385 which resulted in amplification of the entire protein coding regions of the putative M and N genes (5' primer: 5'-GGGGATCCAGAGTTTCAGCGG-3' (SEQ ID NO:30); 3' primer: 5'-GGGAATTCACCACGCATTC-3' (SEQ ID NO:33)). Unique restriction sites (EcoR I and BamH I) at the termini of the PCR products were introduced by conventional methods. A PCR product with the expected size of about 900 bp was obtained

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from each of the virus isolates. Southern blot hybridization was then used to confirm the specificity of the amplified products.

The 32P-labelled cDNA probe from VR 2385 hybridized with the RT-PCR products from each of the above virus The PCR products of the putative M and N genes from each of the PRRSV isolates were purified and cloned into vector pSK+ (Meng et al, J. Vet. Diagn. Invest. 5, 254-258 (1993)). Plasmids containing the full length putative M and N genes were sequenced with an automated DNA Sequencer (obtained from Applied Biosystems, Inc., Foster City California). Three to four cDNA clones from each virus isolate were sequenced with universal and reverse primers, as well as other virus specific sequencing primers (PP288: 5'-GCGGTCTGGATTGACGAC-3' (SEQ ID NO:5) and PP289: 5'-GACTGCTAGGGCTTCTGC-3' (SEQ ID NO:6), each of which is described in application Serial No. 08/131,625, and DP966: 5'-AATGGGGCTTCTCCGG-3' (SEQ ID NO:34)). The sequences were combined and analyzed by the MACVECTOR (International Biotechnologies, Inc.) and GENEWORKS (IntelliGenetics, Inc.) computer programs.

Analysis of the nucleotide sequences encoding the putative M and N proteins of the 5 U.S. PRRSV isolates indicated that, like LV (Meulenberg et al, Virology, 192, 62-72 (1993)) and VR 2385, the putative M and N genes of each of the five additional U.S. isolates overlapped by 8

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base pairs (bp). Figure 17/shows the nucleotide sequence of ORF's 6 and 7 of six U.S. PRRSV isolates and of LV, in which the ISU-12 (VR 2385 and VR 2386) nucleotide sequence (SEQ ID NO:35) is shown first, and in subsequent sequences (SEQ ID NOS:36-41), only those nucleotides which are different are indicated. Start codons are underlined and indicated by (+1>), stop codons are indicated by asterisks (\*), are indicated by (-), and the two larger deletions in the putative N gene are further indicated by (^).

Figures 18(A)-(B) show the alignment of amino acid sequences of the putative M (Fig. 18(A)) and N (Fig. 18(B)) genes of the proposed arterivirus group, performed with a GENEWORKS program (IntelliGenetics, Inc.), using the following parameters (default values): cost to open a gap is 5, cost to lengthen a gap is 25, minimum diagonal length is 4, and maximum diagonal offset is 10. The EAV M gene sequence was omitted because the relatively low sequence identity with PRRSV and LDV requires gaps in the The VR 2385/VR 2386 sequences (SEQ ID NOS:17 alignments. and 19) are shown first, and in subsequent sequences (SEQ ID NOS:43, 45, 47, 49, 51, 24, 53, 55, 57, 59, 61 and 26, respectively), only the differences are indicated. Deletions are indicated by (-), and the two larger deletions in the putative N gene are further indicated by **(^)**.

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Numerous substitutions in the nucleotide sequence were distributed randomly throughout the M and N genes in each of the five isolates, as compared to VR 2385. Most of the substitutions are third base silent mutations when converted to amino acid sequences (see Fig. 18).

Insertions and deletions are found in the nucleotide sequences of the putative M and N genes when comparing the U.S. isolates to LV, but not found among the U.S. isolates (Fig. 17). For example, there are two larger deletions, 15 and 10 nucleotides each, in the putative N gene of the U.S. isolates as compared to the LV N genome (Fig. 17).

The deduced amino acid sequences of the putative M and N genes from the six Iowa strain PRRSV isolates are aligned with the corresponding N sequence of another U.S. isolate, VR 2332 (Murtaugh et al, Proc. Allen D. Leman Swine Conference, Minneapolis, MN, pp. 43-45 (1993)); two European PRRSV isolates, LV (Meulenberg et al, Virology 192, 62-72 (1993)) and PRRSV isolate 10 (PRRSV-10) (Conzelmann et al, Virology, 193, 329-339 (1993)); two LDV strains, LDV-C (Godney et al, Virology, 177, 768-771 (1990)) and LDV-P (Kuo et al, Virus Res., 23, 55-72 (1992)); and EAV (Den Boon et al, J. Virol., 65, 2910-2920 (1991)) (Fig. 18).

The amino acid sequences of the putative N gene are highly conserved among the seven U.S. PRRSV isolates (Fig. 18(B)), and displayed 96-100% amino acid sequence identity

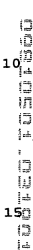
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(Table 1). However, the putative N proteins of the U.S. PRRSV isolates shared only 57-59% amino acid sequence identity with those of the two European isolates (Table 1), suggesting that the U.S. and the European isolates may represent two different genotypes.

The putative M protein of each of the U.S. isolates was also highly conserved, and displayed higher sequence similarity with the M proteins of the two European isolates (Fig. 18(A)), ranging from 78 to 81% amino acid identity (see Table 2 below). The putative N gene of each of the U.S. PRRSV isolates shared 49-50% amino acid sequence identity with that of the LDV strains, whereas the two European PRRSV isolates shared only 40-41% amino acid identity with that of the LDV strains (Table 2).

Two regions of amino acid sequence deletions,

"KKSTAPM" (SEQ ID NO:62) and "ASQG" (SEQ ID NO:63), were

found in the putative N proteins of each of the seven U.S.

PRRSV isolates, as well as the two LDV strains and EAV,

when compared to the two European PRRSV isolates (Fig.

18(B)). These results indicated that the U.S. PRRSV

isolates are more closely related to LDV than are the

European PRRSV isolates, and that PRRSV may have undergone

divergent evolution in the U.S. and in Europe before their

association with PRRS was recognized in swine (Murtaugh,

Proc. Allen D. Leman Swine Conference, Minneapolis, MN, pp.

43-45 (1993)).

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The European isolates may have diverged from LDV for a longer time than the U.S. isolates, and hence may have evolved first. However, the amino acid sequence identity of the putative M gene between U.S. PRRSV isolates and LDV strains was similar to that between the European PRRSV isolates and LDV strains (Table 2). The putative M and N genes of the U.S. and European isolates of PRRSV shared only 15-17% and 22-24% amino acid sequence identity with those of EAV, respectively.

The sequence homology of PRRSV with LDV and EAV suggests that these viruses are closely related and may have evolved from a common ancestor (Plagemann et al, supra; Murtaugh, supra). The high sequence conservation between LDV and PRRSV supported the hypothesis that PRRSV may have evolved from LDV and was rapidly adapted to a new host species (Murtaugh, supra). Asymptomatic LDV infection were found in all strains of mice (Murtaugh, supra; Kuo et al, supra). However, many pig forms are infested with wild rodents (Hooper et al, J. Vet. Diagn. Invest., 6, 13-15 (1994)), so it is possible that PRRSV evolved from LDV-infected mice, and was rapidly adapted to a new host, swine.

The evolutionary relationships of PRRSV with other members of the proposed arterivirus group were determined on the basis of the amino acid sequence of the putative M and N genes. Figure 19 shows a phylogenetic tree of the





Pairwise comparison of the amino acid sequences among the putative nucleocapsid and membrane proteins of the proposed arterivirus group Table 2.

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Virus	VR2385	ISU-22	ISU-55	ISU-79	ISU-1894	ISU-3927	VR2332	LV	PRRSV-10	LDV-P	LDV-C	EAV
VR2385	***	86	96	86	86	96	96	57	57	49	49	22
ISU-22	66	***	86	100	100	86	86	57	57	49	49	23
ISU-55	66	100	***	86	86	. 26	96	59	59	49	49	23
62-USI	86	66	66	***	100	86	86	57	57	49	49	23
ISU-1894	66	100	100	66	***	86	86	57	57	49	49	23
ISU-3927	96	26	26	6	62	***	96	29	59	49	49	23
VR2332	N/A	N/A	N/A	N/A	N/A	N/A	**	57	57	20	49	22
LV	78	62	62	79	79	81	N/A	*	66	41	40	23
PRRSV-10	78	79	79	79	79	81	N/A	91	*	41	6	23
LDV-P	50	51	51	51	51	51	N/A	53	53	*	86	23
LDV-C	49	20	50	20	20	20	N/A	52	52	96	**	24
EAV	16	16	16	91	16	15	N/A	17	17	16	17	*

<sup>b</sup>Nucleocapsid protein comparisons are presented in the upper right half and membrane protein comparisons are presented in the lower left half. Note. \*The values in the table are the percentage identity of amino acid sequences. N/A, not available.

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proposed arterivirus group based on the amino acid sequences of the putative M and N genes of this group. phylogenetic tree for the N gene is essentially the same as that for the M gene. The length of the horizontal lines connecting one sequence to another is proportional to the estimated genetic distance between sequences, as indicated by the numbers given above each line. The UPGMA (unweighted pair group method with arithmetic mean) trees were constructed with a GENEWORKS program (IntelliGenetics, Inc.), which first clusters the two most similar sequences, then the average similarity of these two sequences is clustered with the next most similar sequences or subalignments, and the clustering continued in this manner until all sequences/isolates are located in the tree; both trees are unrooted.

The PRRSV isolates fall into two distinct groups. All the U.S. PRRSV isolates thus far sequenced are closely related and form one group. The two European PRRSV isolates are closely related and form another group. Both the U.S. and European PRRSV isolates are related to LDV strains and are distantly related to EAV (Fig. 19).

The evolution patterns for the putative N and M genes also suggest that PRRSV may be a variant of LDV. For example, the genetic distance of the U.S. PRRSV isolates is slightly closer to LDV than the European PRRSV isolates (Fig. 19), again suggesting that the U.S. and European

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PRRSV may have evolved from LDV at different time periods and existed separately before their association with PRRS was recognized in swine. European PRRSV may have evolved earlier than U.S. PRRSV. It is also possible that the U.S. and European PRRSV could have evolved separately from different LDV variants which existed separately in the U.S. and Europe.

A striking feature of RNA viruses is their rapid evolution, resulting in extensive sequence variation (Koonin et al, Critical Rev. Biochem. Mol. Biol., 28, 375-430 (1993)). Direct evidence for recombination between different positive-strand RNA viruses has been obtained (Lai, Microbiol. Rev., 56, 61-79 (1992)). Western equine encephalitis virus appears to be an evolutionally recent hybrid between Eastern equine encephalitis virus and another alphavirus closely related to Sindbis virus (Hahn et al, Proc. Natl. Acad. Sci. USA, 85, 5997-6001 (1988)). Accordingly, the emergence of PRRSV and its close relatedness to LDV and EAV is not surprising. Although the capsid or nucleocapsid protein has been used for construction of evolutionary trees of many positive-strand RNA viruses, proteins with conserved sequence motifs such as RNA-dependent RNA polymerase, RNA replicase, etc., are typically more suitable for phylogenetic studies (Koonin et al, supra).

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# EXPERIMENT IV:

CLONING AND SEQUENCING OF cDNA CORRESPONDING TO ORF'S 2, 3 AND 4 OF PRRSV VR 2385.

The region including ORF's 2, 3, and 4 of the genome of the porcine reproductive and respiratory syndrome virus (PRRSV) isolate VR 2385 was cloned and analyzed. To clone the cDNA of PRRSV VR 2385, ATCC CRL 11171 cells were infected with the virus at a m.o.i. of 0.1, and total cellular RNA was isolated using an RNA Isolation Kit (Stratagene). The mRNA fraction was purified through a Poly(A) Quick column (Stratagene), and the purified mRNA was used to generate a cDNA library. A cDNA oligo dT library was constructed in Uni-ZAP XR λ vector using a ZAPcDNA synthesis kit (Stratagene), according to the supplier's instructions. Recombinant clones were isolated after screening of the library with an ORF 4 - specific hybridization probe (a 240 b.p. PCR product specific for the 3' end of ORF 4; SEQ ID NO:64). Recombinant pSK + contained PRRSV-specific cDNA was excised in vivo from positive \( \lambda \) plaques according to the manufacturer's instructions.

Several recombinant plasmids with nested set of cDNA inserts with sizes ranging from 2.3 to 3.9 kb were sequenced from the 5' ends of the cloned fragments. The nucleotide sequence of SEQ ID NO:65 was determined on at least two independent cDNA clones and was 1800 nucleotides

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in length (Fig. 21). Computer analysis of the nucleotide and the deduced amino acid sequences was performed using GENEWORKS (IntelliGenetics, Inc.) and MACVECTOR (International Biotechnologies, Inc.) programs.

Three partially overlapping ORF's (ORF 2, ORF 3 and ORF 4) were identified in this region. ORF's 2, 3 and 4 comprised nucleotides 12-779 (SEQ ID NO:66), 635-1396 (SEQ ID NO:68) and 1180-1713 (SEQ ID NO:70), respectively, in the sequenced cDNA fragment.

A comparison of DNA sequences of ORF's 2, 3 and 4 of PRRSV VR 2385 with corresponding ORF's of LV virus (SEQ ID NOS:72, 74 and 76, respectively) is presented in Fig. 22. The level of nucleotide sequence identity (homology) was 65% for ORF 2, 64% for ORF 3 and 66% for ORF 4.

The predicted amino acid sequences encoded by ORF's 2-4 of PRRSV VR 2385 (SEQ ID NOS:67, 69 and 71, respectively) and of LV (SEQ ID NOS:73, 75 and 77, respectively) are shown in Fig 23. A comparison of PRRSV VR 2385 and LV shows a homology level of 58% for the protein encoded by ORF 2, 55% for the protein encoded by ORF 3 and 66% for the protein encoded by ORF 4 (see Fig. 23).

## EXPERIMENT V

An immunoperoxidase method of detecting PRRSV Four 3-week-old colostrum-deprived PRRSV negative animals were inoculated intranasally with 105.8 TCID50 of

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PRRSV U.S. isolate ATCC VR 2386 propagated on ATCC CRL 11171 cells. These pigs were housed on elevated woven-wire decks and fed a commercial milk replacer. Two pigs were necropsied at 4 days post inoculation (DPI) and two at 8 DPI.

At the time of necropsy, the right and left lungs of each pig were separated and inflated via the primary bronchus with 45 ml of one of four fixatives and then immersion fixed for 24 hours. The fixatives used in this experiment included 10% neutral buffered formalin, Bouin's solution, HISTOCHOICE (available from Ambresco, Solon, OH), and a mixture containing 4% formaldehyde and 1% glutaraldehyde (4F:1G). The tissues fixed in Bouin's were rinsed in five 30-minute changes of 70% ethyl alcohol after 4 hours fixation in Bouin's. All the tissues were routinely processed in an automated tissue processor beginning in 70% ethyl alcohol. Tissues were processed to paraffin blocks within 48 hours of the necropsy.

Sections of 3 micron thickness were mounted on poly-1lysine coated glass slides, deparaffinized with two changes
of xylene and rehydrated through graded alcohol baths to
distilled water. Endogenous peroxidase was removed by
three 10-minute changes of 3% hydrogen peroxide. This was
followed by a wash-bottle rinse with 0.05 M TRIS buffer (pH
7.6) followed by a 5-minute TRIS bath. Protease digestion
was performed on all tissue sections except those fixed in

HISTOCHOICE. Digestion was done in 0.05% protease

(Protease XIV, available from Sigma Chem., St. Louis, Mo.)

in TRIS buffer for 2 minutes at 37°C. Digestion was

followed by a TRIS-buffer wash-bottle rinse and then a 5
minute cold TRIS buffer bath. Blocking for 20 minutes was

done with a 5% solution of normal goat serum (available

from Sigma Chem., St. Louis, Mo.).

The primary antibody used was the monoclonal antibody SDOW-17 (obtained from Dr. David Benfield, South Dakota State Univ.), diluted 1:1000 in TRIS/PBS (1 part TRIS:9 parts PBS (0.01 M, pH 7.2)). The monoclonal antibody SDOW-17 recognizes a conserved epitope on the PRRSV nucleocapsid protein (Nelson et al, J. Clin. Microbiol., 31:3184-3189). The tissue sections were flooded with primary antibody and incubated at 4°C for 16 hours in a humidified chamber. primary antibody incubation was then followed by a washbottle rinse with TRIS buffer, a 5-minute TRIS buffer bath, and then a 5-minute TRIS buffer bath containing 1% normal goat serum. The sections were flooded with biotinylated goat anti-mouse antisera (obtained from Dako Corporation, Carpintera, CA) for 30 minutes. The linking antibody incubation was followed by three rinses in TRIS buffer, as was done following primary antibody incubation. sections were then treated with peroxidase-conjugated streptavidin, diluted 1:200 in TRIS/PBS, for 40 minutes, followed by a TRIS buffer wash-bottle rinse and a 5-minute

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TRIS buffer bath. The sections were then incubated with freshly-made 3,3'-diaminobenzidine tetrahydrochloride (DAB, obtained from Vector Laboratories Inc., Burlingame, CA) for 8-10 minutes at room temperature, and then rinsed in a distilled water bath for 5 minutes. Counterstaining was done in hematoxylin (available from Shandon, Inc., Pittsburgh, PA), and the sections were rinsed with Scott's Tap Water (10 g MgSO<sub>4</sub> and 2 g NaHCO<sub>3</sub> in 1 liter ultrapure water), then with distilled water. After dehydration, the sections were covered with mounting media, and then a coverslip was applied.

Two negative controls were included. Substitution of TRIS/PBS buffer in place of the primary antibody was done for one control. The other control was done by substituting uninfected, age-matched, gnotobiotic pig lungs for PRRSV-infected lungs.

Histological changes in infected tissues were characterized by moderate multifocal proliferative interstitial pneumonia with pronounced type 2 pneumocyte hypertrophy and hyperplasia, moderate infiltration of alveolar septa with mononuclear cells, and abundant accumulation of necrotic cell debris and mixed inflammatory cells in the alveolar spaces. No bronchial or bronchiolar epithelial damage was observed. However, there was necrotic cell debris in the smaller airway lumina.

Intense and specific staining in the cytoplasm of infected cells was observed in the formalin- and Bouin's-fixed tissues. Staining was less intense and specific in the 4F:1G-fixed tissues. There was poor staining, poor cellular detail, and moderate background staining in the HISTOCHOICE-fixed tissues. Background staining was negligible with the other fixatives. Cellular detail was superior in the formalin-fixed tissue sections and adequate in the Bouin's- and 4F:1G-fixed tissues.

The labeled antigen was primarily within the cytoplasm of sloughed cells and macrophages in the alveolar spaces (Fig. 24) and within cellular debris in terminal airway lumina (Fig. 25). When compared to sections from the same block stained with hematoxylin and eosin, it was determined that most of the labeled cells were macrophages, and some were likely sloughed pneumocytes. Lesser intensities of staining were observed in mononuclear cells within the alveolar septa and rarely in hypertrophied type 2 pneumocytes.

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Using an immunoperoxidase technique on frozen sections, others were able to detect antigen in epithelial cells of brochioles and alveolar ducts as well as within cells in the alveolar septa and alveolar spaces (Pol et al, "Pathological, ultrastructural, and immunohistochemical changes caused by Lelystad virus in experimentally induced infections of mystery swine disease (synonym: porcine

epidemic abortion and respiratory syndrome (PEARS)), " Vet. Q., 13:137-143). We were unable to detect antigen in brochiolar epithelium using the present immunoperoxidase method.

The present streptavidin-biotin complex (ABC) technique using a PRRSV monoclonal antibody can be modified as needed to identify PRRSV-infected porcine lungs. 10% neutral-buffered formalin and Bouin's solution are acceptable fixatives. Protease digestion enhances the antigen detection without destroying cellular detail. technique is therefore quite useful for the diagnosis of PRRSV-induced pneumonia of pigs, and for detection of PRRSV in lung tissue samples.

#### EXPERIMENT VI

# An immunohistochemical identification of sites of replication of PRRSV

Four three-week-old caesarian-derived, colostrum-Summary: deprived (CDCD) pigs were inoculated intranasally with an isolate of porcine reproductive and respiratory syndrome virus. All inoculated pigs exhibited moderate respiratory disease. Two pigs were necropsied at 4 days post inoculation (PI) and two at 9 days PI. Moderate consolidation of the lungs and severe enlargement of the lymph nodes were noted at necropsy. Moderate perivascular

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lymphomacrophagic myocarditis was observed. Marked lymphoid follicular hyperplasia and necrosis was observed in the tonsil, spleen, and lymph nodes.

Porcine reproduction and respiratory syndrome virus antigen was detected by the present streptavidin-biotin immunoperoxidase method primarily within alveolar macrophages in the lung and in endothelial cells and macrophages in the heart. Macrophages and dendritic-like cells in the lymph nodes, spleen, tonsil, and thymus stained intensively positive for PRRSV nucleocapsid protein antigen as well.

Experimental section: Four pigs were snatched from the birth canal of a sow that was positive for PRRSV antibody by indirect immunofluorescent antibody (IFA) examination of serum. The pigs were taken to a different site, housed on elevated woven-wire decks and raised on commercial milk replacer. These pigs were bled at 0, 7, 14, and 21 days of age and found to be negative for PRRSV antibody by the IFA test. No PRRSV was isolated from the serum of the pigs or sow using MARC-145 cells (available from National Veterinary Services Laboratory, Ames, Iowa).

All four pigs were inoculated intranasally at 3 weeks of age with  $10^{5.8}$  TCID<sub>50</sub> of PRRSV U.S. isolate ATCC VR 2385 propagated on ATCC CRL 11171 cells. Mild-to-moderate respiratory disease was observed from 3-9 days post

inoculation (DPI). Two pigs were necropsied at 4 DPI and two at 9 DPI. At 4 DPI, one pig evidenced 31% and the other 36% tan-colored consolidation of the lungs. At 9 DPI, the remaining two pigs evidenced 37% and 46% consolidation of the lungs, respectively. Lymph nodes were moderately enlarged and edematous.

Lymphoid tissues collected at necropsy included the tonsil, thymus, spleen, tracheobronchial, mediastinal, and medial iliac lymph nodes. Lymphoid tissues were fixed by immersion for 24 hours in 10% neutral buffered formalin, processed routinely in an automated tissue processor, embedded in paraffin, sectioned at 6 microns and stained with hematoxylin and eosin. Additional sections (including the lung tissue sections above) were cut at 3 microns and mounted on poly-L-lysine coated slides for immunohistochemistry.

The immunoperoxidase assay described in Experiment VI above was repeated. Briefly, after endogenous peroxidase was removed with 3% hydrogen peroxide, primary monoclonal antibody ascites fluid diluted 1:1000 in TRIS/PBS was added for 16 hours at 4°C in a humidified chamber. The monoclonal antibody SDOW-17 (obtained from Dr. David Benfield, South Dakota State Univ.), which recognizes a conserved epitope of the PRRSV nucleocapsid protein, was used. Biotinylated goat anti-mouse linking antibody (obtained from Dako Corporation, Carpintera, CA) was added,

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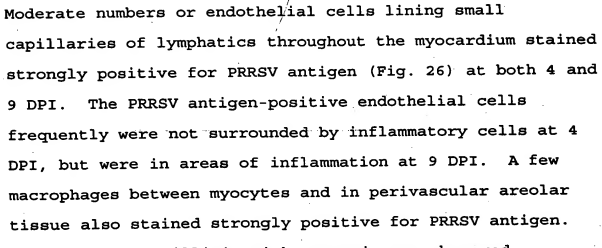
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followed by treatment with peroxidase-conjugated streptavidin (obtained from Zymed Laboratories, South San Francisco, CA) and incubation with 3,3'-diaminobenzidine tetrahydrochloride (obtained from Vector Laboratories Inc., Burlingame, CA). The incubated sample was finally counterstained in hematoxylin.

Microscopic lesions included interstitial pneumonia, myocarditis, tonsillitis, and lymphadenopathy. One section The interstitial of lung from each lobe was examined. pneumonic lesions were characterized by septal infiltration with mononuclear cells, hyperplasia and hypertrophy of type 2 pneumocytes, and accumulation of macrophages and necrotic cell debris in alveolar spaces. These lesions were moderate and multifocal by 4 DPI and severe and diffuse by Bronchi and bronchiolar epithelium was unaffected. PRRSV antigen was readily detected by immunohistochemistry in alveolar macrophages. Large dark-brown PRRSV antigenpositive macrophages were often found in groups of 5-10 cells. A few PRRSV antigen-positive mononuclear cells were observed within the alveolar septa. PRRSV antigen was not detected in any tissues of the negative control pigs.

One section of left and one section of right ventricle were examined. At 4 DPI, there were small, randomly distributed, perivascular foci of lymphocytes and macrophages. There was moderate multifocal perivascular lymphoplasmacytic and histiocytic inflammation by 9 DPI.



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A mild tonsillitis with necrosis was observed.

Necrotic foci of 1-10 cells with pyknosis and karyorrhexis were commonly observed in the center of prominent follicles and less often in the surrounding lymphoreticular tissue.

Large numbers of lymphocytes and macrophages were observed within the crypt epithelium, and moderate amounts of necrotic cell debris were observed in crypts. PRRSV antigen was readily detected within cells in the center of hyperplastic follicles, in the surrounding lymphoreticular tissue, and within cells in the crypt epithelium (Fig. 27). Staining was also present amongst necrotic debris in the crypts. In all these sites, the PRRSV antigen-positive cells resembled macrophages or dendritic-like cells.

Thymic lesions were minimal. There were a few necrotic foci with pyknosis and karyorrhexis in the medulla. These foci tended to involve or be near thymic corpuscles. PRRSV antigen was frequently identified within



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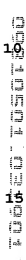
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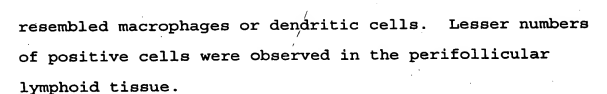
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macrophages near these necrotic areas and less often within large isolated macrophages in the cortex.

Necrotic foci and single necrotic cells were evident with germinal centers of lymphoid nodules and in periarteriolar lymphoid sheaths (PALS) of the spleen. PRRSV antigen positive staining cells were concentrated in the center of lymphoid follicles and scattered throughout The positive cells generally had large oval nuclei and abundant cytoplasm with prominent cytoplasmic projections, compatible with macrophages or dendritic Lesser numbers of positive-staining fusiform-shaped cells in the marginal zone were observed. The size and location of these cells suggests that they are reticular cells.

The predominant lymph node changes were subcapsular edema, foci of necrosis in lymphoid follicles, and the presence of syncytial cells at the border of the central lymphoid tissue with the loose peripheral connective The high endothelial venules were unusually prominent and often swollen. The syncytial cells had 2-10 nuclei with multiple prominent nucleoli and moderate eosinophilic cytoplasm. These cells did not appear to Intense and specific cellular contain PRRSV antigen. cytoplasmic staining was observed in the follicles. positive cells had large nuclei with abundant cytoplasm and prominent cytoplasmic processes (Fig. 27). These cells





The lesion severity and the amount of antigen detected within various tissues was generally similar at 4 and 9 DPI. The gross size of the lymph nodes and the number of syncytial cells in lymph nodes were more prominent at 9 DPI than at 4 DPI. The amount of antigen detected in the heart was also greater at 9 DPI.

Tissues from age-matched uninfected CDCD pigs were used for histologic and immunohistochemical controls.

Other negative controls for immunohistochemistry included using the same protocol less the primary PRRSV antibody on the infected pig tissues. PRRSV antigen was not detected in any of the negative controls.

Conclusions: The immunohistochemical procedure described herein is useful for detecting PRRSV antigen in the lung, heart and lymphoid tissues of PRRSV-infected pigs. Severe interstitial pneumonia and moderate multifocal perivascular lymphohisticcytic myocarditis was observed. Marked lymphoid follicular hyperplasia and necrosis of individual or small clusters of cells in the tonsil, spleen, and lymph nodes was also observed. PRRSV antigen was readily detected in alveolar macrophages in the lung and in endothelial cells and macrophages in the heart.

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Macrophages and dendritic-like cells in tonsil, lymph nodes, thymus, and spleen stained intensely positive for viral antigen as well.

PRRSV may replicate in the tonsil with subsequent viremia and further replication, primarily within macrophages in the respiratory and lymphoid systems of the pig.

#### EXPERIMENT VII

### Diagnosing PRRS:

The present streptavidin-biotin immunoperoxidase test for detection of PRRSV antigen in tissues is quite useful to confirm the presence of active infection. 26 pigs were experimentally inoculated with ATCC VR 2385 PRRSV in accordance with the procedure in Experiments V/VI above. One section of each of the lungs, tonsils, mediastinal lymph nodes, and tracheobronchial lymph nodes from each pig was examined. The virus was detected by the immunoperoxidase assay of Experiment V in 23/26 lungs, 26/26 tonsils, 15/26 mediastinal lymph nodes, and 14/26 tracheobronchial lymph nodes.

The pigs in this experiment were killed over a 28 day period post-inoculation. The virus was detected in at least one tissue in every pig necropsied up to 10 days post inoculation.

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A complete technique for the streptavidin-biotin based immunoperoxidase technique for PRRSV antigen detection in porcine tissues is described in Experiment V infra. Briefly, after endogenous peroxidase removal by 3% hydrogen peroxide and digestion with 0.05% protease (Protease XIV, Sigma Chemical Company, St. Louis, MO), primary monoclonal antibody ascites fluid diluted 1:1000 in TRIS/PBS is added for 16 hours at 4°C in a humidified chamber. The monoclonal antibody used was SDOW-17 (Dr. David Benfield, South Dakota State Univ.), which recognizes a conserved epitope of the PRRSV nucleocapsid protein (Nelson et al, "Differentiation of U.S. and European isolates of porcine reproductive and respiratory syndrome virus by monoclonal antibodies, " J. Clin. Micro., 31:3184-3189 (1993)). Biotinylated goat anti-mouse linking antibody (Dako Corporation, Carpintera, CA) is then contacted with the tissue, followed by treatment with peroxidase-conjugated streptavidin (Zymed Laboratories, South San Francisco, CA), incubation with 3,3'-diaminobenzidine tetrahydrochloride (Vector Laboratories Inc., Burlingame, CA), and finally staining with hematoxylin.

Particularly when combined with one or more additional analytical techniques such as histopathology, virus isolation and/or serology, the present tissue immunoperoxidase antigen detection assay offers a rapid and reliable diagnosis of PRRSV infection.

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# EXPERIMENT VIII

The pathogenicity of PRRSV isolates in 4-8 week old pigs was determined. The isolates were divided into two groups: (1) phenotypes with high virulence (hv) and (2) phenotypes with low virulence (lv) (see Table 3 below). For example, the mean percentage of lung consolidation of groups of pigs inoculated with a PRRSV isolate is shown in Table 4 below. The pathogenicity of a number of PRRSV isolates at 10 DPI is shown in Table 5 below. The results in Table 5 were statistically analyzed to verify the difference between hv and lv phenotypes, as determined by percentage lung consolidation.

Isolates characterized as high virulence produce severe clinical disease with high fever and dyspnea. In general, hv isolates produce severe pneumonia characterized by proliferative interstitial pneumonia with marked type II pneumocyte proliferation, syncytial cell formation, alveolar exudate accumulation, mild septal infiltration with mononuclear cells, encephalitis and myocarditis (designated PRRS-B hereinafter). Isolates characterized as low virulence do not produce significant clinical disease and produce mild pneumonia characterized predominately by interstitial pneumonia with septal infiltration by mononuclear cells, typical of classical PRRS (designated PRRS-A hereinafter).



# Table 3: Characteristics and Pathogenicity of PRRSV Isolates

Virus Isolate	No. of	mRNA 4	Severity of	Microscopic Lesions**		
	Subgenomic mRNAs		gross pneumonia* lesions	Lesion Type in Lung	Heart	Brain
High Virulence (	hv)					
VR 2385	6	Normal	++++	В	++++	++++
VR 2429	8	Normal	++++	В	++++	+++
ISU-28	ND	ND	+++	В	++++	++++
ISU-79	8	Normal	++++	В	+++	+++
ISU-984	ND	ND	+++	В	+++	+++
Low Virulence (	lv)					
ISU-51	··· ND	ND	+	A	+	+
VR 2430	8	Normal	+	A/B	+	+
ISU-95	ND	ND	+	A	+	. +
ISU-1894	6	Normal	+	А/В	+	+_
VR 2431	6	Deletion	+	А/В	-	
Lelystad***	6	Normal	+	A	+/-	+/-

- (-) normal, (+) mild, (++) moderate, (+++) severe, (++++) very severe pneumonia.
- \*\*: PRRSV isolates produce two types of microscopic lung lesions: Type A lesions include interstitial pneumonia with mild septal infiltration with mononuclear cells typical of PRRS as described by Collins et al (1992); Type B lesions include proliferation of type II pneumocytes, and are typical of those described as PIP (Halbur et al 1993).
- \*\*\*: Pol et al, (Vet. Quart., 13:137-143 (1991); Wensvoort et al, Antigenic comparison of Lelystad virus and swine infertility and respiratory syndrome virus. J. Vet. Diagn. Invest., 4:134-138 (1992); Meulenberg et al, Lelystad virus, the causative agent of porcine epidemic abortion and respiratory syndrome (PEARS), is related to LDV and EAV. Virology, 192:62-72 (1993).



## TABLE 4

VIRUS ISOLATE	Mean % Lung Consolidation Score at DPI*				
A	3	10	21	28	
VR-2385	29	77.3	37.3	6.0	
VR-2386pp	20.5	77.5	25.0	0.0	
ISU-22	26.5	64.8	36.5	11.0	
ISU-984	7.25	76.0	21.0	0.5	
ISU-3927	13.5	10.5	0	0.0	
PSP-36	0	0	0	0.0	
UNINOC	0	0	0	0.0	

\*: Score range is from 0-100% consolidation of the lung tissue.

TABLE 5

INOCULUM	NO. PIGS	Mean % Lung Consolidation at 10 DPI <u>+</u> S.D.
Uninfected	10	0 <u>+</u> 0
CRL 11171 Cell Line	10	0 <u>+</u> 0
ISU-51	10	16.7 <u>+</u> 9.0
ISU-55	10	20.8 <u>+</u> 15.1
ISU-1894	10	27.4 ± 11.7
ISU-79	10	51.9 <u>+</u> 13.5
VR-2386pp	10	54.3 <u>+</u> 9.8
ISU-28	10	62.4 <u>+</u> 20.9

\* Pathogenicity of PRRSV isolates ISU-28, VR 2386pp and ISU-79 were not significantly different (p > 0.05) from each other, but were different from that of ISU-51, ISU-55, and ISU-1894 (p < 0.001). All PRRSV isolates were significantly different (p < 0.001) from controls.

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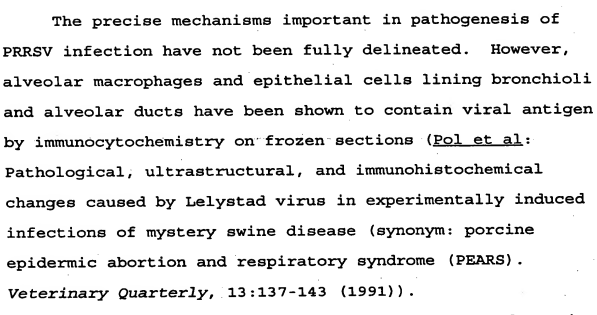
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The present immunocytochemistry test for the detection of PRRSV in formalin-fixed tissues (see Experiment VI supra) shows that PRRSV also replicates in alveolar The extent of virus epithelial cells and macrophages. replication and cell types infected by PRRSV isolates also appears to vary (see Experiment X below).

The role of different genes in virulence and replication is not precisely known. However, ORF's 4 and 5 appear to be important determinants of in vivo virulence and in vitro replication in PRRSV.

The results of cloning and sequencing ORF's 5, 6 and 7 of PRRSV isolate VR 2385 (see Experiment I supra) show that ORF 5 encodes a membrane protein (also see U.S. application Serial No. 08/131,625). A comparison of ORF's 5-7 of VR 2385 with ORF's 5-7 of Lelystad virus shows that ORF 5 is the least-conserved of the three proteins analyzed (see

Table 2 supra), thus indicating that ORF 5 may be important

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EXPERIMENTS IX-XI

PRRSV (ATCC VR 2386) was propagated in vitro in ATCC

CRI. 11171 cells by the method disclosed in Experiment III

in determining virulence.

Based on Northern blot results, ORF 4 of lv isolate VR

2431 appears to have a deletion in mRNA 4 (also see

Experiment V of U.S. application Serial No. 08/131,625).

PRRSV (ATCC VR 2386) was propagated in vitro in ATCC CRL 11171 cells by the method disclosed in Experiment III of U.S. application Serial No. 08/131,625. The PRRSV isolate was biologically cloned by three rounds of plaque purification on CRL 11171 cells and characterized. The plaque-purified isolate (hereinafter "VR 2386pp", which is equivalent to VR 2386, deposited at the ATCC, Rockville Maryland, on October 29, 1992) replicated to about 106-107 TCID<sub>50</sub>/ml at the 11th cell culture passage in CRL 11171 cells. Viral antigens were also detected in the cytoplasm of infected cells using convalescent PRRSV serum. VR 2386pp was shown to be antigenically related to VR 2332 by IFA using polyclonal and monoclonal antibodies to the nucleocapsid protein of VR 2332 (SDOW-17, obtained from Dr. David Benfield, South Dakota State University).

Several other virus isolates (VR 2429 (ISU-22), ISU-28, VR 2428 (ISU-51), VR 2430 (ISU-55), ISU-79, ISU-984, ISU-1894, and VR 2431 (ISU-3927)) were isolated and plaque-purified on CRL 11171 cell line. Virus replication in the

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CRL 11171 cell line varied among PRRSV isolates (see Table 3 below). Isolate VR 2385 and plaque-purified isolates VR 2386pp, VR 2430 and ISU-79 replicated to  $10^{6-7}$  TCID<sub>50</sub>/ml, and thus, have a high replication (hr) phenotype. isolates, such as ISU-984, ISU-1894 replicated to a titer of 10<sup>4-5</sup> TCID<sub>50</sub>/ml, corresponding to a moderate replication (mr) phenotype. Isolates ISU-3927 and ISU-984 replicated very poorly on CRL 11171 cell line and usually yielded a titer of 103 TCID<sub>50</sub>/ml, and thus have a low replication (lr) phenotype.

# EXPERIMENT IX

The pathogenicity of several PRRSV isolates was compared in cesarean-derived colostrum-deprived (CDCD) pigs to determine if there was a correlation between in vitro replication and pathogenicity (also see Experiment V of application Serial No. 08/131,625. Four plaque-purified PRRSV isolates (VR 2386pp, VR 2429, ISU-984, and VR 2431), and one non-plaque-purified isolate (VR 2385) were used to inoculate pigs. An uninoculated group and an uninfected cell culture-inoculated group served as controls. Two pigs from each group were killed at 3, 7, 10, and 21 DPI. pigs were killed at 28 and 36 DPI. Biologically cloned PRRSV isolates VR 2386pp, VR 2429, and ISU-984 induced severe respiratory disease in the 5 week-old CDCD pigs, whereas VR 2431 did not produce any significant disease. Gross lung lesion scores peaked at 10 DPI (see Table 4) and

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ranged from 10.5% consolidation (VR 2431) to 77% consolidation (VR 2385). Lesions were resolved by 36 DPI.

Microscopic lesions included interstitial pneumonia, encephalitis, and myocarditis (Table 3). The lv isolates also caused less severe myocarditis and encephalitis than the hv isolates.

In Figs. 28(A)-(C), photographs of lungs from pigs inoculated with (A) culture fluid from uninfected cell line CRL 11171, (B) culture fluids from cell line infected with lv isolate VR 2431, (C) or culture fluids from cell line infected with hv isolate VR 2386pp. The lung in Fig. 28(B) has very mild pneumonia, whereas lung in Fig. 28(C) has severe consolidation.

#### EXPERIMENT X

An additional experiment was conducted using a larger number of pigs to further examine the pathogenicity of PRRSV isolates and to obtain more statistically significant data. Results are shown in Table 5. Collectively, the results show that PRRSV isolates can be divided into two groups based on pneumopathogenicity. Isolates VR 2385, VR 2429, ISU-28, and ISU-79 have a high virulence (hv) phenotype and produce severe pneumonia. Isolates ISU-51, VR 2430, ISU-1894 and VR 2431 have a low virulence (lv) phenotype (Table 4) and produce low grade pneumonia.



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PRRSV isolates also produce two types of microscopic lesions in lungs. The first type found generally in lv isolates is designated as PRRS-A, and is characterized by interstitial pneumonia with septal infiltration with mononuclear cells typical of PRRS (as described by Collins et al, Isolation of swine infertility and respiratory syndrome virus (isolate ATCC VR-2332) in North America and experimental reproduction of the disease in gnotobiotic J. Vet. Diagn. Invest., 4:117-126 (1992)). second type of lesion, PRRS-B, is found in hv isolates and is characterized as proliferative interstitial pneumonia with marked type II pneumocyte proliferation, alveolar exudation and syncytial cell formation, as described in U.S. application Serial No. 08/131,625 and by Halbur et al, An overview of porcine viral respiratory disease. Proc. Central Veterinary Conference, pp. 50-59 (1993). of PRRS-A and PRRS-B type lesions are shown in Figs. 28(A)-

The immunoperoxidase assay of Experiment V using monoclonal antibodies to PRRSV was used to detect viral antigens in alveolar epithelial cells and macrophages (see Fig. 29(A)). This test is now being routinely used at the Iowa State University Veterinary Diagnostic Laboratory to detect PRRSV antigen in tissues.

(C), in which Fig. 28(A) shows a normal lung, Fig. 28(B)

are the lesions produced by PRRSV type A, and Fig. 28(C)

shows the lesions produced by PRRSV type B.

In Figures 29(A)-(B), immunohistochemical staining with anti-PRRSV monoclonal antibody of lung from a pig infected 9 days previously with VR 2385. A streptavidinbiotin complex (ABC) immunoperoxidase technique coupled with hematoxylin counterstaining were used. Positive staining within the cytoplasm of macrophages and sloughed cells in the alveolar spaces is clearly shown in Fig. 29(A), and within cellular debris in terminal airway lumina in Fig. 29(B).

# EXPERIMENT XI

To determine if there was a correlation between biological phenotypes and genetic changes in PRRSV isolates, Northern blot analyses were performed on 6 PRRSV isolates.

Total intracellular RNA's from the VR 2386pp virusinfected CRL 11171 cells were isolated by the guanidine isothiocyanate method, separated on 1% glyoxal/DMSO agarose gel and blotted onto nylon membranes. A cDNA probe was generated by PCR with a set of primers flanking the extreme 3' terminal region of the viral genome. The probe contained 3' noncoding sequence and most of the ORF-7 sequence (see U.S. application Serial No. 08/131,625).

Northern blot hybridization revealed a nested set of 6 subgenomic mRNA species (Fig. 30). The size of VR 2386pp viral genomic RNA (14.7 kb) and the six subgenomic mRNA's,

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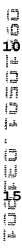


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mRNA 2 (3.3 kb), mRNA 3 (2.8 kb), mRNA 4 (2.3 kb), mRNA 5 (1.9 kb), mRNA 6 (1.4 kb) and mRNA 7 (0.9 kb), resembled those of LV, although there were slight differences in the estimated sizes of the genome and subgenomic mRNA's (Conzelmann et al, Virology, 193, 329-339 (1993), Meulenberg et al, Virology, 192, 62-72 (1993). The mRNA 7 of the VR 2386pp was the most abundant subgenomic mRNA (see Fig. 30 and Experiment I above). The total numbers of subgenomic mRNA's and their relative sizes were also compared. The subgenomic mRNA's of three isolates had 6 subgenomic mRNA's, similar to that described for Lelystad In contrast, three isolates had 8 subgenomic mRNA's virus. (Fig. 30). The exact origin of the two additional species of mRNA's is not known, but they are located between subgenomic mRNA's 3 and 6 and were observed repeatedly in cultures infected at low MOI. Interestingly, an additional subgenomic mRNA has been detected in LDV isolates propagated in macrophage cultures (Kuo et al, 1992). speculate that the additional mRNA's in cells infected with some PRRSV isolates are derived from gene 4 and 5 possibly transcribed from an alternate transcriptional start site. Additional studies are needed to determine the origin of these RNA's and their significance in pathogenesis of PRRSV infections.

Fig. 30 shows Northern blots of PRRSV isolates VR 2386pp (designated as "12"), VR 2429 (ISU-22, designated as





"22"), VR 2430, designated as "55"), ISU-79 (designated as "79"), ISU-1894 (designated as "1894"), and VR 2431, designated as "3927"). This data represents results from four separate Northern blot hybridization experiments. The VR 2386pp isolate (12) was run in one gel, ISU-1894 and VR 2431 were run in a second gel, VR 2430 and ISU-79 were run in a third gel, and ISU-22 was run in a fourth gel. Two additional mRNA's are evident in isolates VR 2429, VR 2430, and ISU-79.

The subgenomic mRNA 4 of VR 2431 (ISU-3927) migrates faster than that of other isolates in Northern blotting, suggesting a deletion. Interestingly, the isolate VR 2431 has lv and lr phenotypes and is the least virulent PRRSV isolate of the Iowa strains described herein. This suggests that gene 4 may be important in virulence and replication. As described above, genes 6 and 7 are less likely to play a role in expression of virulence and replication phenotypes.

In summary, PRRSV isolates vary in pathogenicity and the extent of replication in cell cultures. The number of subgenomic mRNA's and the amount of mRNA's also varies among U.S. PRRSV isolates. More significantly, one of the isolates, VR 2431, which replicates to low titer (lr phenotype) and which is the least virulent isolate (lv phenotype) among the Iowa strain PRRSV isolates described

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herein, appears to have a faster migrating subgenomic mRNA 4, thus suggesting that a deletion exists in its ORF 4.

#### EXPERIMENT XII

COMPARISON OF THE PATHOGENICITY AND ANTIGEN DISTRIBUTION OF TWO U.S. PORCINE REPRODUCTIVE AND RESPIRATORY SYNDROME VIRUS ISOLATES WITH THE LELYSTAD VIRUS

problem on some farms where pig-flow through the unit is appropriate for shedding of the virus from older stock to younger susceptible animals that have lost passive and enterition are protection.

The severity and duration of outbreaks is quite variable. In fact, some herds are devastated by the high production losses (<u>Polson et al.</u>, "Financial Impact of Porcine Epidemic Abortion and Respiratory Syndrome (PEARS)," *Proc. 12th Inter. Pig Vet. Soc.*, p. 132 (1992); <u>Polson et al</u>, "An evaluation of the financial impact of porcine reproductive and respiratory syndrome (PRRS) in nursery pigs," *Proc. 13th Inter. Pig Vet. Soc.*, p. 436

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(1994)), while other herds have no apparent losses due to infection with PRRSV. This may be due to a number of possibilities, including virus strain differences, pig genetic susceptibility differences, environmental or housing differences, or production style (pig flow) of the unit.

This experiment compares the pathogenicity and antigen distribution of two U.S. strains (ISU-12 [VR 2385], ISU-3927 [VR 2431]) and a European strain (Lelystad virus, obtained from the National Veterinary Services Laboratory, P.O. Box 844, Ames, Iowa, 50010) in a common pig model to document similarities and differences that may explain the differences in severity of field outbreaks of PRRSV and help to better understand the pathogenesis of disease induced by PRRSV. (In the following experimental descriptions, "x/y" refers to the number of pigs "x" out of a particular group of pigs having "y" members.)

### Materials and Methods

### Experimental Design:

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One hundred caesarian-derived-colostrum-deprived (CDCD) pigs of 4 weeks of age were randomly divided into 4 large groups of 25 pigs each and assigned to one of four isolated buildings. Within each building, the pigs were further divided into 3 separate rooms (11 pigs, 11 pigs, and 3 pigs per room). Each room within the buildings had

separate, automated ventilation systems. The pigs were housed on raised woven wire decks and fed a complete 18% protein corn and soybean meal based ration. Following challenge with a virus inoculum, the pigs were necropsied as detailed in Table 6 below at 1, 2, 3, 5, 7, 10, 15, 21 and 28 days post inoculation (DPI).

## <u>Virus Inocula Preparation</u>:

Each virus was plaque-purified three times. Challenge doses were  $10^{5.8}$  for VR 2385 and  $10^{5.8}$  for VR 2431. The challenge dose of Lelystad virus was  $10^{5.8}$ .

Pigs were challenged intranasally by sitting them on their buttocks perpendicular to the floor and extending their neck fully back. The inocula was slowly dripped into both nostrils of the pigs, taking approximately 2-3 minutes per pig. Control pigs were given 5 mL of uninfected cell culture media in the same manner.

### Clinical Evaluation

Rectal temperatures were taken and recorded daily from -2 DPI through 10 DPI. A clinical respiratory disease score was given to each pig daily from day 0 to 10 DPI, in accordance with the following 0-6 score range, similar to the respiratory distress analysis described above:

0 = normal

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Table 6: Necropsy Schedule

	T	<u> </u>	The state of the s	
Total	11 11 3	11 11 8	3 3 3	11 11 3
28 DPI	нн	1	<b>п</b> п	пп
21 DPI	11	1	1	ਜ਼ਿਜ
15 DPI	нн	нн	пп -	нн
10 DPI	m m m	ммм	ммм	m m m
7 DPI	пп	нн	нн	ੱਜ ਜ
5 DPI	н н	нн	нн	ਜਜ
3 DPI	ਜਜ	ਜਜ	ਜਜ	пп
2 DPI	нн	нн	нн	11
1 DPI	нн	нн	нн,	нн
Коош	3 2 1	4100	L & 6	10 11 12
Isolate	Lelystad Lelystad Lelystad	VR 2385 VR 2385 VR 2385	Control Control	VR 2431 VR 2431 VR 2431



- 1 = mild dyspnea/and/or tachypnea when stressed
- 2 = mild dyspnea and/or tachypnea when not stressed
- 3 = moderate dyspnea and/or tachypnea when stressed
- 4 = moderate dyspnea and/or tachypnea when not stressed
- 5 = severe dyspnea and/or tachypnea when
   stressed
- 6 = severe dyspnea and/or tachypnea when not stressed

A pig was considered "stressed" by the pig handler after holding the pig under his/her arm and taking the pig's rectal temperature for approximately 30-60 seconds. Other relevant clinical observations like coughing, diarrhea, inappetence or lethargy were noted separately, and are not reflected in the respiratory disease score.

### Pathologic Examination:

Complete necropsies were performed on all pigs.

Macroscopic lung lesions were given a score to estimate the percent consolidation of the lung. Each lung lobe was assigned a number to reflect the approximate volume of entire lung represented by that lobe. Ten (10) possible points were assigned to each of the right anterior lobe, right middle lobe, anterior part of the left anterior lobe, and caudal part of the left anterior lobe of the lung. The accessory lobe was assigned five (5) points. Twenty-seven

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and one-half (27.5) points were assigned to each of the right and left caudal lobes to reach a total of 100 points. Gross lung lesion scores were estimated, and a score was given to reflect the amount of consolidation in each lobe. The total for all the lobes was an estimate of the percent consolidation of the entire lung for each pig.

Sections were taken from all lung lobes, nasal turbinates, cerebrum, thalamus, hypothalamus, pituitary gland, brain stem, choroid plexus, cerebellum, heart, pancreas, ileum, tonsil, mediastinal lymph node, middle iliac lymph node, mesenteric lymph node, thymus, liver, kidney, and adrenal gland for histopathologic examination. Tissues were fixed in 10% neutral-buffered formalin for 1-7 days and routinely processed to paraffin blocks in an automated tissue processor. Sections were cut at 6  $\mu$ m and stained with hematoxylin and eosin.

### Immunohistochemistry:

Immunohistochemical staining was performed as described in Experiment VI above. Sections were cut at 3  $\mu$ m and mounted on poly-L-lysine coated slides. Endogenous peroxidase was removed by three 10-minute changes of 3% hydrogen peroxide. This was followed by a TRIS bath, and then digestion with 0.05% protease (Protease XIV, Sigma Chemical Company, St. Louis, Mo.) in TRIS buffer for 2 minutes at 37°C. After another TRIS buffer bath, blocking

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was done for 20 minutes with a 5% solution of normal goat Primary monoclonal antibody ascites fluid (SDOW-17, obtained from Dr. David Benfield, South Dakota State Univ.) diluted 1:1000 in TRIS/PBS was added for 16 hours at 4°C in a humidified chamber. After primary antibody incubation and a subsequent 5 minute TRIS bath containing 1% normal goat serum, the slides were flooded with biotinylated goat anti-mouse linking antibody (Dako Corporation, Carpintera, CA) for 30 minutes. The sections were washed with TRIS and treated with peroxidase-conjugated streptavidin (Zymed Laboratories, South San Francisco, CA) for 40 minutes, then incubated with 3,3'-diaminobenzidine tetrahydrochloride (Vector Laboratories Inc., Burlingame, CA.) for 8-10 minutes. Sections were then stained with hematoxylin.

Immunohistochemical controls substituted TBS for the primary antibody on all lung and lymphoid tissue sections. The same was done on other sections of other tissues interpreted as possibly positive. Uninfected control pigs also served as negative controls. No staining was detected in any of the control pig tissues. The amount of antigen was estimated according to the following scale: (0) = negative (no positive cells), (1) = isolated or rare positive staining cells (about 1-5 positive cells per histologic section), (2) = a relatively low number of positive cells, yet more abundant than isolated cells (for example, about 10-20 positive cells per histologic

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section), (3) = a moderate number of positive cells (for example, about 40-80 positive cells per histologic section), and (4) = a relatively large number of positive cells (more than about 100 positive cells per histologic section).

#### Virus Isolation:

The same tissues from each of two pigs necropsied from each challenge group were pooled at 1, 2, 3, 5, 7, 14, 21, and 28 DPI. At 10 DPI, nine pigs were necropsied from each group, so three pools of the same tissues from three pigs were made from each challenge group. Serum was also similarly pooled.

#### Results

### Clinical Disease:

The mean clinical respiratory disease score for each group is summarized in Table 7. Control pigs remained normal. Respiratory disease was minimal, and symptoms and histopathology were similar in the groups of pigs infected with Lelystad virus and VR 2431. By 2 DPI, a few pigs in each of these groups demonstrated mild dyspnea and tachypnea after being stressed by handling. From 5-10 DPI, more of the pigs in these groups demonstrated mild respiratory disease, and a couple pigs evidenced moderate, but transient, labored abdominal respiration. By 14 DPI,

Table 7: Mean Clinical Respiratory Disease Score

		_		
10 DPI	0	0.3	0.5	3.0
9 DPI	0.1	0.3	0.5	3.4
8 DPI	0	6.0	0.7	3.3
7 DPI	0.1	1.0	1.3 0.7	3.5
6 DPI	0	0.8	0.3	3.4
5 DPI	0	9.0	9.0	3.2
4 DPI	0	0.5	0.4	2.2
3 DPI	0	0.2	0.2	1.8
2 DPI	0	0.1 0.2	0.3	1.5
1 DPI	0	0.2	0	0.4
0 DPI	0	0	0	0
GROUP	Control	Lelystad	VR 2431	VR 2385

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all pigs in the Lelystad virus (LV) and VR 2431 groups had recovered. Other transient clinical disease noted in a few pigs in these groups included chemosis, reddened conjunctiva, ear drooping, and patchy cyanosis of skin when stressed by handling. Coughing was not observed.

By 2 DPI, the VR 2385-challenged group demonstrated mild respiratory disease without having been stressed. 5 DPI, all of the pigs in this group demonstrated moderate respiratory disease characterized by labored abdominal respiration and dyspnea when stressed. Some of the pigs in this group received respiratory distress scores of 5 or 6 for a 2- to 5-day period, and the mean clinical respiratory disease score peaked at 3.5/6 at 7 DPI. Respiratory disease was characterized by severe tachypnea and labored abdominal respiration, but no coughing was observed. VR 2385 pigs generally were moderately lethargic and anorexic from 4-10 DPI. Other transient clinical signs included chemosis, roughed hair coats, lethargy, and anorexia. It took up to 21 DPI for the majority of the pigs in this group to fully recover.

## Gross Lesions

Table 8 summarizes the estimated percent consolidation of the lungs for pigs in each group. Lung lesions in the Lelystad group and VR 2431 group were similar in type and extent. Lesions were first observed at 5 DPI for both

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groups, and peaked at 15 DPI for the Lelystad challenged group and at 7 DPI for VR 2431 challenged group. Individual scores ranged from 0-31 percent consolidation for the Lelystad group and 0-27 percent for the VR 2431 The mean estimated percent consolidation of the lung for the nine pigs necropsied at 10 DPI was 6.8 percent for Lelystad virus challenged pigs and 9.7 percent for the VR 2431 challenged pigs. The lesions were predominately in the cranial, middle and accessory lobes and in the ventromedial portion of the diaphragmatic lobes. The consolidation was characterized by multifocal, tan-mottled areas with irregular, indistinct borders.

Table 8: Estimated Percent Consolidation of Lungs (0-100%)

GROUP	1	2	3	5	7	10	15	21	28
	DPI	DPI	DPI	DPI	DPI	DPI	DPI	DPI	DPI
Control	0 0 0	0	0	0	0	0	0	0	0
Lelystad		0	0	4.8	2.3	6.8	8.8	1.8	0
VR 2431		0	0	2.5	8.5	9.7	7.5	0	0
VR 2385		4.3	10.5	15.3	46.5	54.2	12.5	6.0	0

Gross lymphoid lesions were more common than lung lesions with both VR 2431 and LV. Lymphadenopathy was consistently observed in the mediastinal and middle iliac lymph nodes. These lymph nodes were tan in color, and from 5-28 DPI, were enlarged to 2-10 times their normal size. There often was at least one 1-5 mm fluid-filled cyst in

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each of these lymph nodes.  $\Big/$  No other gross lesions were

observed in the LV or VR 2431 groups.

The VR 2385 group had considerably more severe lung consolidation. The distribution of lung consolidation was similar to pigs infected with VR 2431 and LV, but either the entire cranioventral lobes or large coalescing portions of the cranial, middle, accessory and ventromedial diaphragmatic lobes were consolidated. There was no pleuritis and no grossly visible pus in airways. Estimated percent consolidation of the lung 7-10 DPI ranged from 28% to 71%. The estimated mean score of the nine pigs necropsied at 10 DPI was 54.2% consolidation.

Lymphoid lesions in the VR 2385 group were generally similar to those observed in the other groups.

Additionally, lymph nodes along the thoracic aorta and in the cervical region were often 2-5 times the normal size. Spleens were also slightly enlarged and meaty in texture.

Several pigs in the VR 2385 group had moderately enlarged and rounded hearts with 10-30 mL of clear fluid in the pericardial space. Some of these pigs also had 50-200 mL of similar fluid in the abdominal cavity. There was no visible exudate or fibrin in the fluid.

## Microscopic Lesions:

<u>Heart</u>: Control pigs necropsied up to 10 DPI had no evidence of myocardial inflammation. Several pigs

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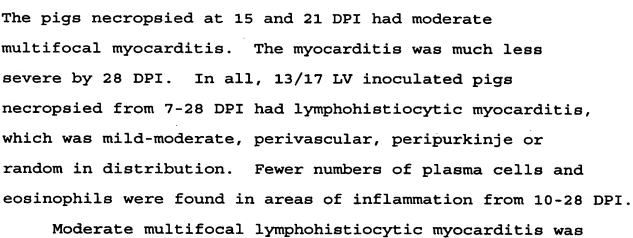
throughout the study had randomly distributed discrete foci of hematopoietic cells in the endocardium and myocardium. These hematopoietic cells (i) were observed in clumps of 10-30 cells, (ii) ranged in size from 8-20 microns, and (iii) had large round-oval, dark staining nuclei with dense, clumped chromatin, multiple small nucleoli and scant amphophilic cytoplasm. At 10 DPI, 2/9 control pigs had mild multifocal perivascular lymphohisticcytic myocarditis. This was also observed in 1/2 pigs necropsied at 15 and 21 DPI, respectively.

VR 2431 inoculated pigs also had evidence of myocardial extramedullary hematopoiesis, similar to the controls. Myocarditis was first observed at 7 DPI, and was seen in 16/18 pigs necropsied from 7-28 DPI. myocarditis was mild, multifocal, usually perivascular and peripurkinje, and lymphohistiocytic. Inflammation was consistently found in the endocardium, often around or involving purkinje fibers. Inflammation in the epicardium and myocardium was most consistently either around vessels or randomly distributed between muscle fibers. Myocardial degeneration, necrosis, or fibrosis was not evident. numbers of eosinophils were observed in the perivascular infiltrates in a 4/9 pigs at 9 DPI.

In the LV inoculated pigs, mild multifocal extramedullary hematopoiesis was evident in most pigs up to 7 DPI. Mild myocarditis was first observed at 2 DPI and

was inconsistent and mild in pigs posted from 3-10 DPI.

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Moderate multifocal lymphohistic myocarditis was observed beginning at 10 DPI in all of the VR 2385 inoculated pigs. Severe myocarditis was observed in 2/9 pigs killed at 10 DPI and in 1/2 pigs killed at each of 15, 21, and 28 DPI, respectively. The more severe cases were characterized by multifocal-to-diffuse, lymphoplasmacytic and histiocytic infiltrates that were most intense in the perivascular, peripurkinje, and endocardial regions.

Lesser numbers of eosinophils and unidentifiable pyknotic cells were also observed in association with the inflammation. Myocardial degeneration, necrosis and fibrosis were not evident.

Lung: Very mild lung lesions were observed in 2/25 of the control pigs. One pig necropsied at 5 DPI had mild multifocal septal thickening with lymphocytes, macrophages, and neutrophils. At 10 DPI, one pig had mild peribronchiolar and perivascular lymphohistiocytic cuffing

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and a mild increased number of macrophages and neutrophils in the alveolar spaces.

In the VR 2431 inoculated pigs, microscopic lung lesions were first detected at 2 DPI and were present in 20/25 of the pigs. All pigs necropsied on or after 7 DPI had microscopic lung lesions. The lesions, when present, were multifocal, mild (12/25) to moderate (8/25), generally most severe at 10 DPI and nearly resolved at 28 DPI. multifocal interstitial pneumonia was characterized by three primary changes: septal thickening with mononuclear cells, type 2 pneumocyte hypertrophy and hyperplasia, and accumulation of normal and necrotic macrophages in alveolar spaces. These changes were present throughout the 28-day period. Mild-to-moderate peribronchiolar and perivascular lymphohistiocytic cuffing was observed in most pigs examined at 10-15 DPI but had apparently resolved by 28 Lung lesions were seldom observed in sections taken from the caudal lung lobe.

The LV inoculated pigs had microscopic lung lesions very similar to those of VR 2431 in distribution, type, and severity. Microscopic lung lesions were observed in 21/25 of the LV pigs. Lesions were first observed at 2 DPI and persisted throughout the 28 day period. The most severe lesions were seen in a few of the pigs necropsied at 10 DPI and in most of those necropsied at 15 and 21 DPI. The interstitial pneumonia was characterized mainly by septal

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thickening with mononuclear cells, peribronchiolar and perivascular lymphohisticcytic cuffing, and accumulation of macrophages and necrotic debris in alveolar spaces. Type 2 pneumocyte hyperplasia and hypertrophy was less consistent and less severe than that observed in the VR 2431 inoculated pigs. Lung lesions were seldom seen in sections taken from the caudal lung lobe.

Every pig that was inoculated with VR 2385 and necropsied on or after 5 DPI had moderate-to-severe interstitial pneumonia. Mild multifocal lesions were observed at 2 DPI. The lesions became moderate and multifocal by 5 DPI, severe and diffuse from 7-10 DPI, and still moderate but patchy at 21 and 28 DPI. The interstitial pneumonia at all stages was also characterized by three primary changes (septal thickening with mononuclear cells, type 2 pneumocyte hypertrophy and hyperplasia, and accumulation of normal and necrotic macrophages in alveolar spaces). Of these three changes, the pneumocyte hypertrophy was most prominent and characteristic of VR 2385 inoculation. Peribronchiolar and perivascular lymphomacrophagic cuffing was mild by 5 DPI, moderate by 10 DPI, and nearly resolved by 28 DPI.

#### Immunohistochemistry

Both adrenal glands were examined from all pigs.

Adrenal gland lesions were not observed in any of the



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control, VR 2431 or LV inoculated pigs. In the VR 2385 inoculated pigs, 9/25 pigs had mild multifocal lymphoplasmacytic and histiocytic adrenalitis.

Inflammation was usually observed in the medulla. Pyknotic cells and karryhectic debris were also observed amongst the inflammatory cells. Lymphoplasmacytic vasculitis and neuritis were also observed in the adrenal artery and nerve, respectively, in 3/28 of the VR 2385 inoculated pigs.

Nasal turbinate lesions were similar in type but differed in severity and frequency in the 4 groups of pigs. A low number (5/25) of the control and LV (5/25) inoculated pigs had mild rhinitis, observed at 10-21 DPI. The rhinitis was characterized by patchy dysplasia of the epithelium, with loss of cilia and mild multifocal subepithelial lymphohisticcytic and suppurative inflammation, with slight edema and congestion.

More of the VR 2431 inoculated pigs (17/25) had rhinitis. Lesions were mild at 5 DPI but moderate by 10 DPI. Epithelial dysplasia with intercellular edema, a blebbed or "tombstone" appearance of swollen superficial epithelial cells becoming pyknotic and apparently sloughing into the nasal cavity, and complete or partial loss of cilia on large patches of epithelium were observed. There was moderate diffuse subepithelial edema, dilated and congested veins, and multifocal infiltrates of lymphocytes,

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plasma cells, macrophages and neutrophils. The inflammation was most intense near the locations where the ducts of submucosal mucous glands extended to the surface. Leukocytic exocytosis, especially of neutrophils, were frequently observed in dysplastic surface epithelium and along mucous ducts. By 21 DPI, the lesions had become mild, and were resolved by 28 DPI.

Rhinitis was first observed at 5 DPI in the VR 2385 inoculated pigs. A total of 20/25 pigs, and all 17 pigs necropsied on or after 7 DPI, had rhinitis similar to that observed in the ISU-3927 group, except that the lesion persisted throughout the 28 day period.

Tables 9, 10, and 11 summarize and compare the number of different tissues in which PRRSV antigen was detected for each of the challenge groups. No antigen was detected in the control pigs. Table 12 summarizes the estimated amount of antigen in some of the tissues that were tested.

#### Virus isolation

Virus isolation from various tissues is summarized in Table 13, where "Lg" refers to lungs, "LN" refers to lymph nodes, "Ht" refers to the heart, "Ser" refers to serum, "Tons" refers to tonsils, "Spln" refers to the spleen, "SI" refers to small intestine, and "Brn" refers to the brain.

Table 9: Immunohistochemistry for VR 2385

Tissue	1 DPI	2 DPI	3. DPI	5 DPI	7 DPI	10 DPI	15 DPI	12 Taa	28 DPI	Total
Lung TBLN Med LN Iliac LN Tonsil Thymus Spleen	0/2 1/2 0/2 2/2 0/2	2/22	2000000	2000 20/00 2	2/2 2/2 2/2 0/2 0/2	9/8 3/9 3/9 3/9	2/2 0/2 0/2 0/2 0/2 0/2	2/2 1/2 0/2 0/2 0/2 1/2	0/2 0/2 0/2 0/3	22/25 13/25 14/25 14/25 25/25 9/25
sod #	2/2	2/2	2/2	2/2	2/2	6/6	2/2	2/2	2/2	25/25

Table 10: Immunohistochemistry for VR 2431

Total	14/25 8/25 10/25 8/25 21/25 6/25 1/25	25/25
28 DPI	2/2 0/2 0/2 0/2 1/2	2/2
21 DPI	0/22000/200000/2000/2000/2000/2000/2000/2000/2000/2000/2000/2000/2000/20000/2	2/2
15 DPI	//2222 //000/000	2/2
10 DPI	7/9 11/9 11/9 0/9 0/9	6/6
7 DPI	0/1700	2/2
5 DPI	1/2 2/2 2/2 2/2 1/2 0/2	2/2
3 DPI	0/2 1/2 2/2 1/2 0/2	2/2
2 DPI	1/2 2/2 2/2 0/2 0/2	2/2
DPI	0/2	1/2
Tissue	Lung TBLN Med LN Iliac LN Tonsil Thymus Spleen	sod #

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Table 11: Immunohistochemistry for Lelystad virus

<u>م</u> ا	ស្ត្រស្ត្រស្ត្	52
Total	14/25 10/25 10/25 23/25 7/25	25/25
28 DPI	7555555	2/2
21 DPI	2222222 000000000000000000000000000000	2/2
15 DPI	000000	2/2
10 DPI	5/9 2/9 0/9 4/9	8/9
7 DPI	1/2 1/2 1/2 2/2 0/2	2/2
Ida	1/2 0/2 0/2 2/2 0/2	2/2
3 DPI	1/2 1/2 2/2 2/2 0/2	2/2
2 DPI	1/2 1/2 1/2 2/2 0/2 1/2	2/2
1 DPI	0/1 1/2 1/3 1/3 1/3 1/3 1/3 1/3 1/3 1/3 1/3 1/3	2/2
Tissue	Lung TBLN Med LN Iliac LN Tonsil Thymus Spleen	god #

# Serology

All pigs challenged with LV virus were negative prechallenge and remained <1:20 through 7 DPI. By 10 DPI, 6/9 of the pigs necropsied were seropositive with titers ranging from 1:20 to 1:1280. Only 2/10 pigs had titers >1:20 (both were 1:1280). By 15 DPI, all pigs were positive and 5/6 were >1:320. By 21 DPI, titers of 1:1280 or 1:5120 were most common. The VR 2431 antibody titers were similar to those levels seen with the LV virus. With VR 2385, however, 9/9 were positive by 10 DPI and 7/9 were ≥1:320. No PRRSV serum antibody was detected in control pigs.

#### Discussion

This Experiment clearly demonstrates differences in pathogenicity between PRRSV isolates, differences in PRRSV antigen distribution, and differences in the amount of PRRSV antigen in selected tissues. The low virulence Iowa strain isolate VR 2431 and the low virulence Lelystad virus were similar in these criteria. The Iowa strain VR 2385 isolate was considerably more virulent, and PRRSV antigen was detected in more tissues and in greater amounts as compared to LV and VR 2431.

The pattern of antigen distribution over time (Table 12) suggests that when pigs are infected oronasally, initial and continual replication of the virus may be in

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Table 12: Mean score for intensity/amount of PRRSV antigen detected by immunohistochemistry

													L					
			æ	VR 2385					YR.	VR 2431					Lelystad Virus	ad Vi	rus	
CrVn M. Lung Lu	ΣĀ	Mid Lung	TBLN	Med	lliac LN	Tonsil	CrVn Lung	Mid Lung	TBLN	Med	lliac LN	Tonsil	CrVn lung	Mid Lung	TBLN	Med LN	lliac LN	Tonsil
·°	0		1.5	o	0.5	1.0	0.5	0	0	0	0	0.5	0	0	0.5	0.5	0	1.0
0.5	"	1.0	2.0	1.5	2.0	1.5	0.5	0	2.0	1.0	2.5	0.5	0.5	0.5	1.0	1.0	1.0	1.0
2.0	'''	2.5	3.0	3.0	3.0	3.0	0	0	1.0	1.5	2.5	0.5	0.5	0.5	1.0	1.5	2.0	1.0
2.0	1	2.0	3.0	3.0	2.5	3.0	0.5	н	2.0	2.0	2.0	1.0	1.0	0.5	0	0.5	<b>o</b> /	1.0
2.5	1/	1.5	1.0	1.5	2.0	1.0	0	7	1.0	1.0	0.5	0.5	1.0	0	1.0	0.5	0.5	1.0
2.0		1.6	0.5	9.6	0.7	1.2	1.1	6.0	0.1	0.1	0.1	1.1	0.3	0.4	0.6	0.2	0	9.0
1.0	1	۰	0	0	0	1.0	2.0	5.0	0	0	0	1.0	0.5	0.5	0	0	0	1.0
2.0		0.5	0.5	0	0	2.5	0	ò	0	0	0	1.0	1.0	0	0	0.5	0	1.5
1.0		٥	0	1	.0	1.5	1.3	0	0	1.3	0	2.0	0.5	0	0	0.5	0	1.0
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Antigen amount was estimated and scored as follows: (0) = negative, (1) = isolated or rare positive staining cells, (2) = low number of positive cells, (3) moderate number of positive cells, and (4) = large number of positive cells.

CrVn = Cranioventral lung lobe; Mid = middle lung lobe; TBLN = tracheobronchial lymph node; Med LN = mediastinal lymph node.

Table 13: Virus isolation

			12	VR 2385							Λ	VR 2431						1	elyst	Lelystad Virus	ug		
ž.	E	出	Ser	Tons	Spln	SI	Brn	Гg	LIN	Ht	Ser	Ser Tons	Spln	SI	Brn	Lg	IN	Ht	Ser	Tons	Spln SI		Bri
1	+		+			+		+		ı	+	+	,	+		•	ı		+	+		•	b
	+	+	+	+	+	+	+	+	+	+	+	+	+	+		+	+	+	+	+	+	+	<b>)</b> '
	+	+	+	+	+	+	+	+	+	+	+	÷	+	+	•	+	+	+	+	+	+	+	ı
	+	+	+	+	+	+	+	+	+	+	+	+	+		+	+	+	ı	+	+	+	+	1
	+	+	+	+	+	+	+	+	+	+	+	<b>+</b>	+	•	+	+	+	+	<b>+</b> .	+	+	+	•
	+	+	+	+	+	+		+	+	+	+	+	+	•	+	+	+	+	+	+	+	+	1
	+	•	+	+	+	+	•	+	+	+	+	+	+	•		+	+	+	+	+	+	+	ı
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	+	+	+	+	+	+	•	+	•	1	+	+				+	•	+	+	+	,	1	•
	+	•	+	+	·		Ī	. +	1	+	+	+				+		+	+	+			



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the tonsil and upper respiratory tract lymphoid tissues, with subsequent viremia by 24 hours PI. A small amount of antigen is detected in the lung by 24 hours PI and peaks by 5-7 DPI, but persists there for up to 28 days. Antigen is present in lymphoid tissues generally from 2-21 DPI.

Antigen is detected primarily within the macrophages and dendritic-like cells in lung, lymph nodes, tonsil, thymus and spleen.

#### EXPERIMENT XIII

COMPARATIVE PATHOGENICITY OF NINE U.S. PRRSV ISOLATES IN A 5 WEEK OLD CDCD PIG MODEL

Part (A) of this experiment demonstrates a consistent model to study PRRSV-induced respiratory and systemic disease in piglets (e.g., about 5 weeks old) and to characterize gross and microscopic lesions associated with the course of PRRSV-induced disease. Part (B) of this experiment uses the model to statistically compare the virulence of PRRSV isolates from herds with differing disease severity, and to specifically determine if these differences may be due to virus virulence characteristics.

#### Materials and Methods

#### Source of PRRSV isolates:

Live pigs or fresh tissues were received from 61 herds over a 3-year period from 1991-1993. All cases were

submitted for etiologic diagnosis of respiratory disease in pigs from 1-16 weeks of age. Some of the herds had concurrent reproductive failure, and some did not. nine selected herds differed in size, production style, age of diseased pigs, time since initial disease was observed, and severity of the current disease outbreak. The clinical information from the selected farms is summarized in Table 14.

Table 14: PRRSV Herd Profiles

Isolate	Herd Size	Production Style	Age of Disease	Type of Disease
VR 2385	180 sows	F-Fin/CF	ALL	severe PRRS
ISU-79	40 sows	F-Fin/AIAO	ALL	severe PRRS
ISU-28	150 sows	F-Fin/CF	ALL	severe PRRS
ISU-1894	600 sows	F-FRP/CF	3-8 weeks	severe resp.
VR 2428	900 sows	F-FRP/AIAO	3-8 weeks	severe resp.
VR 2429	100 sows	F-Fin/CF	1-8 weeks	moderate resp.
ISU-984	600 sows	F-FRP/AIAO	3-6 weeks	moderate resp.
VR 2430	150 sows	F-Fin/CF	3-6 weeks	mild resp.
VR 2431	60 sows	F-Fin/AIAO	1-4 weeks	mild resp.

F-Fin = Farrow-to-Finish

F-FRP = Farrow-to-Feeder Pig

CF = Continuous Flow AIAO = All-in-All-out

## Inocula preparation

PRRSV isolates were plaque purified 3 times in accordance with the procedure described in Experiment I, section (I)(A) above.

#### Experimental pigs:



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Four-week-old caesarean-derived-colostrum-deprived (CDCD) pigs were initially fed a commercial 22% protein pig starter containing spray-dried plasma protein for 7 days, then were switched to a second stage 18% protein cornsoybean meal based ration for the duration of the experiment. Pigs were housed in 10 feet x 12 feet concrete-floored, individually power-ventilated rooms.

## Part (A): CDCD pig model:

Ninety-eight 4-week-old CDCD pigs were randomly divided into 7 rooms of 14 pigs each. The rooms were randomly assigned one of seven treatments as shown in Table The treatment consisted of intranasal inoculation of 105.7 TCID<sub>50</sub> of a PRRSV isolate (selected from plaquepurified PRRSV isolates VR 2385, VR 2428 [ISU-22], VR 2431 or ISU-984, unplaque-purified isolate ISU-12 [VR 2386]), intranasal inoculation of uninfected cell culture and media, or no treatment. Two pigs from each group were necropsied at DPI 3, 7, 20 and 21, and 3 pigs were necropsied from each group at DPI 28 and 36. temperatures were recorded daily from DPI -2 though DPI +14. A clinical respiratory disease score was given from DPI -2 through DPI 14. Scores range from 0-6, in accordance with the respiratory distress scale recited in Experiment XII. A piglet was considered "stressed" by the pig handler when holding the pig under his/her arm and

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taking the rectal temperature for approximately 30-60 seconds. Other relevant clinical observations (e.g., coughing, diarrhea, inappetence or lethargy) were noted separately as observed. Additional clinical observations had no impact on the clinical respiratory score. Weights were recorded an DPI 0, 7, 14, 21 and 28.

Table 15: Part (A) Experimental Design

Inoculum	3 DPI	7 DPI	10 DPI	21 DPI	28 DPI	36 DPI	Total Pigs
VR 2385	2	2	2	2	3	3	14
ISU-984	2	2	2	2	3	3	14
VR 2428	2	2	2	2	3	3	14
VR 2431	2	2	2	2	3	3	14
VR 2386	2	2	2	2	3	3	14
Uninoculated Control	2	2	2	2	3	3	14
PSP-36 Cell Culture	2	2	2	2	3	3	14

## Part (B): Comparative Pathogenicity:

Results from Part (A) established that gross lung lesions were most severe at 10 DPI for 4 of 5 PRRSV isolates. Part (B) was designed to collect and compare data from a larger number of pigs necropsied at 10 DPI. In this experiment, 105 4-week-old crossbred CDCD pigs were randomly divided into seven rooms, each with 15 pigs. Each room was randomly assigned a/treatment. Treatments

consisted of intranasal challenge with 10<sup>5.8</sup> TCID<sub>50</sub> of one of six plaque-purified PRRSV isolates (VR 2429 [ISU-51], ISU-79, VR 2430 [ISU-55], ISU-1894, ISU-28 or VR 2385) or PSP-36 uninfected cell culture and media. Ten pigs from each group were necropsied at 10 DPI, and 5 pigs from each group were necropsied at 28 DPI. Rectal temperatures were recorded from -2 DPI to +10 DPI, and weights were recorded at 0, 10 and 28 DPI. Clinical respiratory disease scores and other clinical signs were recorded as in Part (A) above.

#### Serology:

Part (A): Pigs were bled at 0, 10 and 28 DPI. The
presence of PRRSV serum antibody was detected by the
immunofluorescent antibody technique (IFA) as described by
Benfield et al (J. Vet. Diagn. Invest., 4:127-133 (1992)).

Part (B): Pigs were bled at 0, 3, 10, 16 and 28 DPI
and tested by the IFA procedure of Part (A) for the
presence of PRRSV serum antibody.

## Virus Isolation:

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Virus isolation was attempted from lung homogenates of all pigs killed at 3, 7, 10, 21 and 28 DPI (Part (A)).

Virus isolation was also attempted from lung and from serum of all pigs separately in two-pig pools using CRL 11171 (PSP 36) cells (Part (B)).

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## Gross Pathology:

Complete necropsies were performed on all pigs. All organ systems were examined. An estimated percent consolidation of the lung of each pig was calculated based on the scoring system described in Experiment XII above, in which each lung lobe was assigned a number to reflect the approximate volume of entire lung represented by that lobe. Other lesions were noted accordingly.

## Microscopic Pathology:

Sections were taken from all lung lobes described above, as well as from nasal turbinates, cerebrum, thalamus, hypothalamus, pituitary gland, brain stem, choroid plexus, cerebellum, heart, pancreas, ileum, tonsil, mediastinal lymph node, middle iliac lymph node, mesenteric lymph node, thymus, liver, kidney, and adrenal gland for histopathologic examination. Tissues were fixed in 10% neutral buffered formalin for 1-7 days and routinely processed to paraffin blocks in an automated tissue processor. Sections were cut at 6  $\mu$ m and stained with hematoxylin and eosin. Lesions in several tissues were graded in accordance with the following scale: (-) = normal, (+) = mild, (++) = moderate, (+++) = severe, and (++++) = very severe (see Table 19).



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#### Results

# Clinical disease - Part (A), CDCD pig model:

VR 2385 challenged pigs demonstrated the most severe clinical respiratory disease, with scores above 2.5/6.0 on 7-9 DPI (Table 16). The onset of respiratory disease was noted on 3 DPI, and symptoms and lesions continued through Respiratory disease was characterized by labored and accentuated abdominal respirations and tachypnea. There was no coughing. The pigs became lethargic by 3 DPI, were anorexic by 5 DPI, and did not return to full feed and activity until after 14 DPI. Eyelid edema was noted in two pigs on 6 and 7 DPI.

VR 2428-challenged pigs had a later onset of respiratory disease (5 DPI), but severe respiratory disease occurred more quickly and for a longer duration than in ISU-12-inoculated pigs. VR 2428 produced respiratory scores greater than 3.0/6.0 on 7-13 DPI. The pigs were off feed and lethargic at 6-14 DPI. No other clinical signs were noted.

ISU-984-challenged pigs produced moderate-to-severe respiratory disease with gradual onset starting at 4 DPI. The pigs were scored 2-2.5/6.0 for respiratory disease from 7-10 DPI, and greater than 3.0/6.0 with a few scores of 4-5/6.0 on 11-14 DPI. Other clinical signs included lethargy, eyelid edema, and blotchy-purple transient discoloration of skin.

VR 2431-challenged pigs produced mild respiratory disease. Disease onset occurred at 5 DPI with the most severe respiratory clinical disease scores between 2 and 2.5/6.0 in some pigs at 7-8 DPI. The pigs appeared considerably better by 10 DPI and were completely normal by 14 DPI. Lethargy and anorexia were observed on 7-8 DPI.

Mean rectal temperatures were greater than 104°F for all challenged groups by 7 DPI, and remained above 104°F until after 10 DPI. This coincided with the period of most severe clinical respiratory disease. The control pigs remained clinically normal throughout the experiment.

## Clinical disease - Part (B), Comparative pathogenicity:

Clinical respiratory disease scores and rectal temperatures are summarized in Table 17. VR 2429 produced very mild respiratory disease and the pigs appeared near normal through 10 DPI. VR 2430 induced mild dyspnea and tachypnea from 4-10 DPI, as well as lethargy and anorexia from 4-6 DPI. At 5-8 DPI, ISU-1894 produced moderate respiratory disease of short duration, and the pigs were generally recovered by 10 DPI. ISU-1894-inoculated pigs were also transiently lethargic and anorexic from 4-7 DPI. ISU-79 induced severe respiratory disease with labored respirations of increased frequency, accompanied by lethargy and anorexia from 4 DPI to 15 DPI. ISU-12 induced moderate tachypnea and dyspnea of long duration (4-28 DPI).



These pigs were also moderately lethargic and mildly anorexic over that time period.

Pigs in three groups (ISU-12, ISU-79, ISU-28) frequently exhibited transient, blue-purple discoloration of the skin when stressed by handling. ISU-28 produced severe respiratory disease similar to ISU-79, but had a later onset (at 7 DPI) and only a 5-day duration. Controls remained normal through 10 DPI.

## Gross lesions - Part (A), CDCD pig model:

Gross lung lesions were scored and estimated as percent lung consolidation. Results are summarized in Table 16. The degree of consolidation ranged from 7.3% (ISU-984) to 29% (VR 2386) at 3 DPI, 20% (VR 2431) to 56.3% (VR 2386) at 7 DPI, 10.5% (VR 2431) to 77.5% (VR 2385) at 10 DPI, 0% (VR 2431) to 37.3% at 21 DPI, and 0% (VR 2431, VR 2385) to 11% (VR 2428) at 28 DPI. No grossly detectable lesions remained in any group at 36 DPI. No gross lung lesions were observed at any time in the control group.

The affected lung lobes were primarily in the anterior, middle, accessory, and ventromedial portion of the caudal lobes. The consolidated areas were not well demarcated. These areas were multifocal within in each lobe and had irregular and indistinct borders, giving the affected lobes a tan-mottled appearance.

Table 16: Part (A) Mean Gross Lung Consolidation

		3 [	PI	7 E	PI	10 1	OPI	21 [	PI	28 1	PI
	Isolate	Clin. Score	Gross Lung								
	VR 2386	0.5	29	3.1	56.3	3.5	77.3	2.0	37.3	0.5	6.0
	VR 2385	0.5	20.5	2.3	35.5	2.0	77.5	0.5	25.0	0	0.0
	VR 24289	0	26.5	2.4	35.0	3.5	64.8	2.0	36.5	2.5	11.0
	ISU-984	0.5	7.3	2.3	21.8	3.5	76.0	2.0	21.0	0	0.0
FF 11.11 I	VR 2431	0	13.5	2.3	20.0	1.5	10.5	0	0	0.	0.0
11 11 11.	PSP-36	0	0	0	О	0	0	0	0 .	. <b>o</b>	0.0
R. H 2211	Uninoc.	0	0	0	0	0	0	0	0	0	0.0

## Gross lesions - Part (B), Comparative pathogenicity:

Gross lung lesions were estimated by percent lung consolidation, and are shown in Table 18.

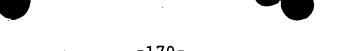
## Microscopic lesions - Part (A), CDCD pig model:

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Results are shown in Table 19. VR 2385, VR 2386, VR 2428 and ISU-984 all induced similar microscopic lung lesions. They produced moderate-severe interstitial pneumonia, characterized by: (i) type II pneumocyte proliferation, (ii) septal thickening with mononuclear cells, and (iii) accumulation of mixed alveolar exudate. VR 2431 induced only mild interstitial pneumonia with

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septal thickening by mononuclear cells. Myocarditis was observed only in the VR 2386 inoculated pigs.

## Virus Isolation - Part (A), CDCD pig model:

PRRSV was recovered from the lungs of all 11 pigs inoculated with VR 2386, from 9 of 11 pigs inoculated with VR 2385, from 6 of 11 pigs inoculated with ISU-984, from 9 of 11 pigs inoculated with VR 2431, from 0 of 11 pigs inoculated with cell culture controls, and from 0 of 11 uninoculated control pigs up to 28 DPI.

### Serology - Part (A), CDCD pig model:

All of the PRRSV inoculated pigs had detectable PRRSV antibody titer of  $\geq$  640 by 10 DPI. None of the control pigs had detectable PRRSV antibody. Most of the PRRSVinoculated pigs had titers of  $\geq$  2560 by 28 DPI.

#### <u>Serology - Part (B), Comparative pathogenicity:</u>

All of the PRRSV-inoculated pigs had PRRSV antibody titers of  $\geq$  64 by 10 DPI. Control pigs did not have detectable PRRSV antibody.

#### Discussion

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The 5-week-old CDCD pigs inoculated intranasally with 10<sup>5.8</sup> TCID<sub>50</sub> of PRRSV provide an excellent model to study and compare PRRSV-induced respiratory and systemic disease.







Significant differences (p < .05) were observed in the pneumopathogenicity data reported in Table 18. Based on the results herein and in Experiment XI above, the isolates could be grouped into high and low virulence groups as follows:

high virulence: VR 2385, VR 2386, VR 2429 (ISU-22),
ISU-28, ISU-984, ISU-79

low virulence: VR 2431, VR 2428 (ISU-51), VR 2430, ISU-1894, LV

A PRRSV isolate may be considered to be a "high virulence" phenotype if it results in one or more of the following:

- (a) a mean gross lung consolidation at 10 DPI of at least 30%, and preferably, at least 40%;
- (b) moderate-to-very severe type II pneumocyte hypertrophy and hyperplasia, moderate-to-very severe interstitial thickening, moderate-to-very severe alveolar exudate, and the presence of syncytia; or
- (c) a mean respiratory distress score of at least 2.0 at some point in time from 10-21 DPI.

Where an isolate does not meet any of the above criteria, it may be considered a "low virulence" phenotype.





Obviously, numerous modifications and variations of the present invention are possible in light of the above teachings. It is therefore to be understood that within the scope of the appended claims, the invention may be practiced otherwise than as specifically described herein.

Table 17: Part (B) Mean Respiratory Distress Scores and Mean Rectal Temperature (°F)

T		T	1	<del></del>				
	28 DPI	103.8	104.2	104.1	103.9	103.8	103.8	103.9
	21 DPI	103.5	103.6	104.6	104.4	103.5	104.2	103.8
Temperature	15 DPI	103.1	104.5	103.5	103.9	103.4	103.7	104.0
Rectal Temp	10 DPI	103.7	103.2	103.8	103.3	103.7	103.5	104.8
Mean Rec	7 DPI	103.3	104.2	104.1	104.3	104.6	103.9	104.0
	5 DPI	102.6	103.7	103.7	104.4	104.9	104.3	104.2
·	3 DPI	102.7	102.6	102.8	102.7	103.6	102.2	102.6
ıre	28 DPI	0	0	0	0	1.0	0.8	0
ss Score	21 DPI	0	0.2	0	0	0.5	2.4	0
istre	15 DPI	0	0	0	0.5	1.5	1.0	0
ory D	10 DPI	0	0.2	1.5	1.1	2.9	1.4	3.1
spirat	7 DPI	0	0.7	0.8	1.5	3.8	1.4	1.3
Mean Respiratory Distress	5 DPI	0	0.1	1.1	2.5	3.5	1.5	1.0
Me	3 DPI	0	0	0	0	0	0.2	
	Isolate	PSP-36	VR 242,98	VR 2430	ISU-1894	ISU-79	VR 2385	ISU-28





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Table 18: Part (B), Mean Gross Lung Consolidation and Standard Deviation

Inocula	Number of Pigs	Mean gross lung score 10 DPI	SD
PSP-36	10	0.0	0.0
ISU-28	10	62.4	20.9
VR 2385	10	54.3	9.8
ISU-79	10	51.9	13.5
ISU-1894	10	27.4	11.7
VR 2430	10	20.8	15.1
VR 2429%	10	16.7	9.0

VR 2429 10 16.7 9.0

Table 19: Experiment XIII, part (A), CDCD pig model: Microscopic Lesion
Summary at 10 DPI

Lesion	VR 2386	VR 2385	VR 2428	ISU-984	VR 2431	PSP-36 control
Type II pneumocyte proliferation	++++	+++	+++	+++	+	-
Syncytia	++	++	++	++	-	-
Interstitial thickening	++++	+++	+++	+++	+	-
alveolar exudate	+++	+++	+++	+++	+	<u>-</u>
myocarditis	+	-	-	-	-	-
encephalitis	+	- ,	-	-	-	-

#### SEQUENCE LISTING

#### (1) GENERAL INFORMATION:

- (i) APPLICANT: PAUL, PREM S. MENG, XIANG-JIN HALBUR, PATRICK G. MOROZOV, IGOR LUM, MELISSA A.
- (ii) TITLE OF INVENTION: A POLYNUCLEIC ACID ISOLATED FROM A PORCINE REPRODUCTIVE AND RESPIRATORY SYNDROME VIRUS (PRRSV), A PROTEIN ENCODED BY THE POLYNUCLEIC ACID, A VACCINE PREPARED FROM OR CONTAINING THE POLYNUCLEIC ACID OR PROTEIN,
- (iii) NUMBER OF SEQUENCES: 77
  - (iv) CORRESPONDENCE ADDRESS:
    - (A) ADDRESSEE: OBLON, SPIVAK, McCLELLAND, MAIER & NEUSTADT, P.C.
    - (B) STREET: 1755 S. Jefferson Davis Highway, Suite 400
    - (C) CITY: Arlington
    - (D) STATE: Virginia
    - (E) COUNTRY: U.S.A.
    - (F) ZIP: 22202
    - (v) COMPUTER READABLE FORM:
      - (A) MEDIUM TYPE: Floppy disk

      - (B) COMPUTER: IBM PC compatible (C) OPERATING SYSTEM: PC-DOS/MS-DOS
      - (D) SOFTWARE: PatentIn Release #1.0, Version #1.25
  - (vi) CURRENT APPLICATION DATA:
    - (A) APPLICATION NUMBER: US
    - (B) FILING DATE:
    - (C) CLASSIFICATION:
- (vii) PRIOR APPLICATION DATA:
  - (A) APPLICATION NUMBER: US 08/131,625
  - (B) FILING DATE: 05-OCT-1993
- (viii) ATTORNEY/AGENT INFORMATION:
  - (A) NAME: Lavalleye, Jean-Paul M.P.
  - (B) REGISTRATION NUMBER: 31,451
  - (C) REFERENCE/DOCKET NUMBER: 4625-021-55X CIP
  - (ix) TELECOMMUNICATION INFORMATION:
    - (A) TELEPHONE: (703) 413-3000
    - (B) TELEFAX: (703) 413-2220
    - (C) TELEX: 248855 OPAT UR

	(2)	INFO	DRMATION FOR SEQ ID NO:1:		
		(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 22 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: unknown  (D) TOPOLOGY: linear		
		(ii)	MOLECULE TYPE: DNA (genomic)	,	
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F. 2 2		(ii)	) MOLECULE TYPE: DNA (genomic)		
		(xi)	) SEQUENCE DESCRIPTION: SEQ ID NO:2:		
	CCC	CATTT	TCC CTCTAGCGAC TG		22
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		(i)	) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 20 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: unknown  (D) TOPOLOGY: linear		
		(ii)	) MOLECULE TYPE: DNA (genomic)		
		(xi)	) SEQUENCE DESCRIPTION: SEQ ID NO:3:		
	~~~	00003	33.G. G1MG13.GG1G		

	(2)	INFO	RMATION FOR SEQ ID NO:4:	
	·	(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 20 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: unknown  (D) TOPOLOGY: linear	
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14		(ii)	MOLECULE TYPE: DNA (genomic)	
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	(2)			
		(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 20 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: unknown	
			(D) TOPOLOGY: linear	
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		(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:6:	
	GAC	TGCTA	AGG GCTTCTGCAC	2
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(B) TYPE: nucleic acid

		(C) STRANDEDNESS: unknown (D) TOPOLOGY: linear			
•	(ii)	MOLECULE TYPE: DNA (genomic)			
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(2	) INFO	RMATION FOR SEQ ID NO:8:			
The Car and	(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 19 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: unknown  (D) TOPOLOGY: linear			
that north that the that that	(ii)	MOLECULE TYPE: DNA (genomic)			
i ===	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:8:			
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	) INFO	RMATION FOR SEQ ID NO:9:			
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	(ii)	MOLECULE TYPE: DNA (genomic)			
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	(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 19 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: unknown  (D) TOPOLOGY: linear			

		(ii)	MOLECULE TYPE: DNA (genomic)	
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	TTG	ACGAG	GA CTTCGGCTG	19
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		(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 20 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: unknown  (D) TOPOLOGY: linear	· -
		(ii) <sub>.</sub>	MOLECULE TYPE: DNA (genomic)	
4 22	:	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:11:	
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l÷.		(ii)	MOLECULE TYPE: DNA (genomic)	
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		(ii)	MOLECULE TYPE: cDNA	



## (vi) ORIGINAL SOURCE:

- (A) ORGANISM: porcine reproductive and respiratory syndrome virus
- (B) STRAIN: Iowa
- (C) INDIVIDUAL ISOLATE: ISU-12 (VR 2385/VR 2386)

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:

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		GGCAGTGGAG	TGTTTTGTCA	TTTTTCCTGT	GTTGACTCAC	ATTGTCTCTT	660
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	TCATCATAGA	GAAAAGGGGC	AAAGTTGAGG	TCGAAGGTCA	CCTGATCGAC	CTCAAAAGAG	960
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	GTCACCTATT	CAATTAGGGC	GACCGTGTGG	GGGTAAGATT	TAATTGGCGA	GAACCACACG	2040
=	GCCGAAATTA	ААААААААА	AA				2062

(2) INFORMATION FOR SEQ ID NO:14:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 603 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: unknown
  - (D) TOPOLOGY: unknown
- (ii) MOLECULE TYPE: cDNA
- (vi) ORIGINAL SOURCE:
  - (A) ORGANISM: porcine reproductive and respiratory syndrome virus
  - (B) STRAIN: Iowa
  - (C) INDIVIDUAL ISOLATE: ISU-12 (VR 2385/VR 2386)
- (ix) FEATURE:

ıŌ

- (A) NAME/KEY: CDS
- (B) LOCATION: 1..600
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:14:

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																Asn	
		AAC Asn														TGT Cys	144
		CTG Leu 50														GTG Val	192
		TGT Cys														GGT Gly 80	240
		CTC Leu														Val	288
		ACC Thr													Met	TAC Tyr	336
		GTC Val							Ile					Arg		GCG Ala	384
		AAT Asn 130											Tyr			TTT Phe	432
•		CTG Leu										Arg				ATC Ile 160	480
		GAG Glu									Gly					Leu	528
	Lys		Val	Val 180	Leu	Asp	Gly	Ser	Ala 185	Ala					Arg	GTT , Val	576
		GCG Ala		Gln					)								603



# (2) INFORMATION FOR SEQ ID NO:15:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 200 amino acids
  - (B) TYPE: amino acid
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:15:

Met Leu Gly Lys Cys Leu Thr Ala Gly Cys Cys Ser Gln Leu Leu Phe 1 5 10 15

Leu Trp Cys Ile Val Pro Ser Cys Phe Val Ala Leu Val Ser Ala Asn 20 25 30

Gly Asn Ser Gly Ser Asn Leu Gln Leu Ile Tyr Asn Leu Thr Leu Cys 35 40 45

Glu Leu Asn Gly Thr Asp Trp Leu Ala Asn Lys Phe Asp Trp Ala Val 50 55 60

Glu Cys Phe Val Ile Phe Pro Val Leu Thr His Ile Val Ser Tyr Gly 65 70 75 80

Ala Leu Thr Thr Ser His Phe Leu Asp Thr Val Gly Leu Val Thr Val 85 90 95

Ser Thr Ala Gly Phe Val His Gly Arg Tyr Val Leu Ser Ser Met Tyr 100 105 110

Ala Val Cys Ala Leu Ala Ala Leu Ile Cys Phe Val Ile Arg Leu Ala 115 120 125

Lys Asn Cys Met Ser Trp Arg Tyr Ser Cys Thr Arg Tyr Thr Asn Phe 130 135 140

Leu Leu Asp Thr Lys Gly Arg Leu Tyr Arg Trp Arg Ser Pro Val Ile 145 150 155 160

Ile Glu Lys Arg Gly Lys Val Glu Val Glu Gly His Leu Ile Asp Leu 165 170 175

Lys Arg Val Val Leu Asp Gly Ser Ala Ala Thr Pro Val Thr Arg Val 180 185 190

Ser Ala Glu Gln Trp Ser Arg Pro 195 200

M 11 Mars 11

	(2)	INFO	RMAT	ION	FOR	SEQ	ID N	10:16	5: /								•
		(i)	(A (B (C	) LE ) TY !) SI	E CH INGTH IPE: IRANI IPOLO	: 52 nucl EDNE	5 ba eic SS:	se p acid unkr	oairs 1	<b>;</b>							
		(ii)	MOL	ECUI	E TY	PE:	CDNA	7									
		(vi)	( <i>P</i>	) OF	vii RAIN	SM: rus I: Ic	poro wa		repr E: IS							y syn	drome
F. 45 17 17	*	(ix)	FEA	TURE		ŒY:	CDS			·.					,		
		(xi)	SEC	OUEN	CE DI	ESCR	IPTIC	ON: S	SEQ :	ID NO	0:16	:					
		GAG Glu														Gln	4.8
		GTG Val													Ile		96
4		CTA Leu												His		TTG Leu	144
		TTC Phe 50														CAC His	192
	TTT Phe 65	Gln	AGT Ser	ACA Thr	AAT Asn	AAG Lys 70	GTC Val	GCG Ala	CTC Leu	ACT Thr	ATG Met 75	Gly	GCA Ala	GTA Val	GTT Val	GCA Ala 80	24
		CTT Leu									Thr					Thr	28
	TCC	AGA	TGC	CGT	TTG	TGC	TTG	CTA	GGC	CGC	AAG	TAC	ATT	CTG	GCC	CCT	33

Ser Arg Cys Arg Leu Cys Leu Leu Gly Arg Lys Tyr Ile Leu Ala Pro

105

110

100

The trans over over one. The start

										CAT (					AT Asn		384
GAT Asp	AAC Asn 130	CAC His	GCA Ala	TTT Phe	GTC Val	GTC Val 135	CGG Arg	CGT Arg	CCC Pro	GGC '	rcc z Ser 140	ACT I	ACG ( Thr	TC I Val	AAC Asn		432
										GTG ' Val 155							480
										AAA Lys							522
[]TAA																	525
(2)  ≟	INF	ORMA"	rion	FOR	SEQ	ID :	NO:1	7:									
		(i) s	(A) (B)	LEI TYI	NGTH PE: 8	: 17 amin	ERIS' 4 am o ac line	ino a id		s							
j																	
	(:	ii) ľ	MOLE	CULE	TYP	E: p	rote	in			*						
l						_			Q ID	NO:	17:						
- E	(: Glu	xi) :	SEQUI	ENCE	DES	CRIP	TION	: SE		Asp	•	Thr	Ala	Pro 15	Gln		
13 13 15 14 14 15 15 16 16 16 16 16 16 16 16 16 16 16 16 16	(: Glu	xi) : Ser	SEQUI Ser	ENCE Leu 5	DES	CRIP Asp	TION Phe	: SE	His 10	Asp	Ser			15		* 1	
U G Met E 1	Glu Val	xi) : Ser Leu	SEQUI Ser Leu 20	ENCE Leu 5 Ala	DES Asp Phe	CRIP Asp Ser	TION Phe	: SECCYS Thr 25	His 10 Tyr	Asp	Ser	Val	Met 30	15 Ile	Tyr		
Met Lys	(: Glu Val Leu	Ser Leu Lys 35 Leu	SEQUI Ser Leu 20 Val	Leu 5 Ala Ser	DES Asp Phe Arg	CRIP Asp Ser	TION Phe Ile Arg 40	: SE Cys Thr 25 Leu	His 10 Tyr	Asp	Ser Pro	Val Leu 45 Thr	Met 30 His	15 Ile Leu	Tyr Leu		
□ □ Met □ 1 Lys Ala	Glu Val Leu Phe 50	Ser Leu Lys 35 Leu	SEQUI Ser Leu 20 Val	Leu 5 Ala Ser Cys	DES Asp Phe Arg	CRIP Asp Ser Gly Phe 55	TION Phe Ile Arg 40	: SE Cys Thr 25 Leu Phe	His 10 Tyr Leu Gly	Asp Thr	Pro Leu Met 60 Gly	Val Leu 45 Thr	Met 30 His	Ile Leu Val	Tyr Leu His		
Lys Ala Val	Glu Val Leu Phe 50	Ser  Leu  Lys 35  Leu  Ser	SEQUI Ser Leu 20 Val Asn	Leu 5 Ala Ser Cys Asn	DESCASP Phe Arg Ala Lys 70 Tyr	CRIP Asp Ser Gly Phe 55	TION Phe Ile Arg 40 Thr	: SECCYS Thr 25 Leu Phe	His 10 Tyr Leu Gly	Asp Thr Gly Tyr Met 75	Pro Leu Met 60 Gly	Val Leu 45 Thr	Met 30 His Phe Val	15 Ile Leu Val	Tyr Leu His Ala 80		

Ala His His Val Glu Ser Ala Ala Gly Phe His Pro Ile Ala Ala Asn

			115			•		120	,'				125					
1	Asp	Asn 130	His	Ala	Phe	Val	Val 135	Arg	Arg	Pro	Gly	Ser 140	Thr	Thr	Val	Asn		
	Gly 145	Thr	Leu	Val	Pro	Gly 150	Leu	Lys	Ser	Leu	Val 155	Leu	Gly	Gly	Arg	Lys 160		
•	Ala	Val	Lys	Gln	Gly 165	Val	Val	Asn	Leu	Val 170	Lys	Tyr	Ala	Lys				
	(2)	INF	ORMAI	CION	FOR	SEQ	ID 1	10:18	В:									
int tind the		(i	(E	A) LE 3) TY C) ST	ENGTI (PE : [RAN]	HARACH: 3' nuc. DEDNI OGY:	72 ba leic ESS:	ase j acie unk	pair: d nown	<b>3</b>								
		(ii	) MOI	LECUI	LE T	YPE:	CDN	A										
m this and the		(vi	(1	A) OI B) S:	RGAN: vi: ral:	OURC ISM: rus N: I IDUA	por owa									y syr	ndrome	<b>.</b>
			(1	A) NA B) L	AME/	KEY: ION:	1	369										
		-	) SE							•			•	٠				
	Met		AAT Asn			Gly					Arg					Gly		48
			GTC Val							Leu					. Ala	CAC His	٠.	96
	CAA	AAC	CAG	TCC	AGA	GGC	AAG	GGA	CCG	GGA	AAG	AAA	AAT	AAG	AAG	AAA		144

Gln Asn Gln Ser Arg Gly Lys Gly Pro Gly Lys Lys Asn Lys Lys

AAC CCG GAG AAG CCC CAT TTC CCT CTA GCG ACT GAA GAT GAT GTC AGA

35

50

Asn Pro Glu Lys Pro His Phe Pro Leu Ala Thr Glu Asp Asp Val Arg

	CAT His 65	CAC His	TTT Phe	ACC Thr	CCT Pro	AGT Ser 70	GAG Glu	CGT Arg	CAA Gln	TTG Leu	TGT Cys 75	CTG Leu	TCG Ser	TCA . Ser	ATC Ile	CAG Gln 80	240
	ACC Thr	GCC Ala	TTT Phe	AAT Asn	CAA Gln 85	GGC Gly	GCT Ala	GGG Gly	ACT Thr	TGC Cys 90	ACC Thr	CTG Leu	TCA Ser	GAT Asp	TCA Ser 95	GGG Gly	288
	AGG Arg	ATA Ile	AGT Ser	TAC Tyr 100	ACT Thr	GTG Val	GAG Glu	TTT Phe	AGT Ser 105	TTG Leu	CCT Pro	ACG Thr	CAT His	CAT His 110	ACT Thr	GTG Val	336
	CGC Arg	CTG Leu	ATC Ile 115	CGC Arg	GTC Val	ACA Thr	GCA Ala	TCA Ser 120	CCC Pro	TCA Ser	GCA Ala	TGA					372
	] [(2)	TNF	אמרר	rion	FOD	SEO.	ו חד		۵.	•							·
į	ĝ				•												•
, m	h I		(1) :		LEI	NGTH	: 123	3 am:	ino		s						
100			٠					o ac: linea		•							
į	<u>.</u>	(:	ii) ľ	MOLE	CULE	TYPI	E: 101	rote:	in								
i	No.	_		SEQUI						Q ID	NO:	19:					
de mente sentes sentes	Met	(3	xi) s	SEQUI	ENCE	DES	CRIP'	rion	: SE	_			Lys	Gly	Asp 15		
A 16 18 18 18 18 18 18 18 18 18 18 18 18 18	Met	(z Pro	xi) {	SEQUI Asn	ENCE Thr 5	DES	CRIP:	FION Gln	: SE	Lys 10	Arg	Lys			15		
A 16 18 18 18 18 18 18 18 18 18 18 18 18 18	Met Met 1 Gln	Pro	xi) 8 Asn Val	SEQUI Asn Asn	Thr 5 Gln	DESO Gly Leu	CRIP'	FION Gln Gln	: SE Gln Met 25	Lys 10 Leu	Arg Gly	Lys	Ile	Ile 30	15 Ala	His	
A 16 18 18 18 18 18 18 18 18 18 18 18 18 18	Met 1 Gln	Pro Pro Asn	xi) s Asn Val Gln 35	Asn Asn 20 Ser	Thr 5 Gln Arg	DESC Gly Leu Gly	CRIP Lya Cya Lya	Gln Gln Gly 40	: SE Gln Met 25 Pro	Lys 10 Leu Gly	Arg Gly Lys	Lys	Ile Asn 45	Ile 30 Lys	15 Ala Lys	His	
A 16 18 18 18 18 18 18 18 18 18 18 18 18 18	Met Met I Gln Gln Asn	Pro Pro Asn Pro 50	Asn Val Gln 35 Glu	Asn Asn 20 Ser Lys	Thr 5 Gln Arg	DESO Gly Leu Gly	CRIPT Lys Cys Lys Phe 55	Gln Gln Gly 40 Pro	: SE Gln Met 25 Pro	Lys 10 Leu Gly Ala	Arg Gly Lys Thr	Lys Lys Glu 60	Ile Asn 45 Asp	Ile 30 Lys Asp	15 Ala Lys Val	His Lys	
A 16 18 18 18 18 18 18 18 18 18 18 18 18 18	Met Met 1 Gln Gln Asn His	Pro Pro Asn Pro 50	Xi) S Asn Val Gln 35 Glu Phe	Asn Asn 20 Ser Lys	Thr 5 Gln Arg Pro	DESC Gly Leu Gly His Ser 70	Lys Lys Lys Phe 55	Gln Gly 40 Pro	: SE Gln Met 25 Pro Leu Gln	Lys 10 Leu Gly Ala Leu	Arg Gly Lys Thr Cys 75	Lys Lys Glu 60 Leu	Ile Asn 45 Asp	Ile 30 Lys Asp	15 Ala Lys Val	His Lys Arg Gln 80	
A 16 18 18 18 18 18 18 18 18 18 18 18 18 18	Met Met Gln Gln Asn His 65	Pro Pro Asn Pro 50 His	Asn Val Gln 35 Glu Phe	Asn Asn 20 Ser Lys	Thr 5 Gln Arg Pro Pro Gln 85	DESC Gly Leu Gly His Ser 70 Gly	Lys Cys Lys Phe 55 Glu	Gln Gly 40 Pro Arg Gly	Gln Met 25 Pro Leu Gln Thr	Lys 10 Leu Gly Ala Leu Cys 90	Arg Gly Lys Thr Cys 75	Lys Lys Glu 60 Leu Leu	Ile Asn 45 Asp Ser	Ile 30 Lys Asp Ser	15 Ala Lys Val Ile Ser	His Lys Arg Gln 80	

	(2)	INFO	RMAI	'ION	FOR	SEQ	ID N	iO:20	): /		,							
		(i)	( <u>P</u> (E	L) LE 3) TY C) SI	CE CHENGTH PE: TRANI	i: 60 nucl EDNE	06 ba Leic ESS:	acio unkr	pairs 1	3								
		(i,i)	MOI	ECUI	LE TY	PE:	CDNA	Ā						٠			•	
-		(vi)	(2	A) OF	AL SO RGANI Vii VDIVI	SM: cus	por		. –			e and	d res	spira	atory	y synd	drome	
		(ix)		4) N2	E: AME/I DCATI			503					·					•
H THE		(xi)	SEÇ	QUEN	CE DI	ESCR:	IPTIC	ON: S	SEQ :	ID NO	0:20	:						
					CAC His 5													48
					TTT Phe												·	96
÷ ~~					GAC Asp												1	.44
					AAT Asn								His				1	.92
					TTT Phe							Thr				TCA Ser 80	. 2	240
					ACA Thr 85						Asp					Gly	2	288
										Gly					Cys	AGC Ser	3	336

													TTT G Phe 125					384
													ACC C					432
												Arg	TGG I					480
											Val		GGC ? Gly					528
· ID										Val			CAA (					576
	Arg				GAG Glu				Ala									606
i		INF	ORMA'	rion	FOR	SEQ	ID	NO:2	1:					,				
		·	(i) :	(A (B	ENCE ) LEI ) TY ) TO	NGTH PE:	: 20 amin	l am o ac	ino id		ls							
ļ.		(	ii)	MOLE	CULE	TYP	E: p	rote	in									
	٠	(:	xi)	SEQU	ENCE	DES	CRIP	TION	: SE	Q II	NO:	21:			ŕ			
	Met 1	Arg	Cys	Ser	His 5	Lys	Leu	Gly	Arg	Phe 10		Thr	Pro	His	Ser 15			
	Phe	Trp	Trp	Leu 20		Leu	Leu	Сув	Thr 25		Leu	ı Ser	Trp	Ser 30		Ala		
-	Asp	Gly	Asn 35	Gly	Asp	Ser	Ser	Thr 40		Glr	туг	: Ile	Tyr 45	Asn	Leu	Thr	•	
	Ile	Сув 50		Leu	Asn	Gly	Thr 55		Tr	Let	ı Ser	Ser 60		Phe	Gl	7 Trp	٠	
	Ala 65		Glu	Thr	Phe	Val		туг	Pro	Va.	L Ala 75		r His	Ile	Let	Ser 80		

	Leu	Gly	Phe	Leu	Thr 85	Thr	Ser	His	Phe	Phe 90	Asp	Ala	Leu	Gly	Leu 95	Gly		
	Ala	Val	Ser	Thr 100	Ala	Gly	Phe	Val	Gly 105	Gly	Arg	Tyr	Val	Leu 110	Cys	Ser		
	Val	Tyr	Gly 115		Cys	Ala	Phe	Ala 120	Ala	Phe	Val	Cys	Phe 125	Val	Ile	Arg		,
	Ala	Ala 130	Lys	Asn	Cys	Met	Ala 135	Cys	Arg	Tyr	Ala	Arg 140	Thr	Arg	Phe	Thr		
	Asn 145	Phe	Ile	Val	Asp	Asp 150	Arg	Gly	Arg	Val	His 155	Arg	Trp	Lys	Ser	Pro 160		
( = F		Val	Val	Glu	Lys 165	Leu	Gly	Lys	Ala	Glu 170	Val	Asp	Gly	Asn	Leu 175	Val		
	Thr	Ile	Lys	His 180	Val	Val	Leu	Glu	Gly 185	Val	Lys	Ala	Gln	Pro 190	Leu	Thr		
	Arg	Thr	Ser 195	Ala	Glu	Gln	Trp	Glu 200	Ala									
4	(2)	INF	ORMA'	TION	FOR	SEQ	ID I	NO:2	2:									
FINE FEET			) SE( () ()	QUENCA) LI B) T C) S	CE CI ENGTI YPE: IRANI	HARA	CTER 54 ba leic ESS:	ISTIC ase j acic unki	CS: pair d	S								
		(ii)	) MO	LECU	LE T	YPE:	CDN	A									÷ .	
		(vi	()	IGINZ A) O B) S C) I	RGAN: vi: TRAI	ISM: rus N: I	por owa							_		y syn	drome	
		(xi	) SE	QUEN	CE D	ESCR	IPTI	ON:	SEQ	ID N	0:22	:						
	TGG	GCTG	GCA '	TTCT	rgag(	GC A	rccci	AGTG:	r TTC	BAAT'	rgga	AGAZ	ATGCC	TG C	TGAA	TGGC	A	60
	CTG	ATTG	ACA '	TTGT	GCCT	CT A	AGTC	ACCT	A TTC	CAAT	ragg	GCGA	ACCGI	GT G	GGGG	TAAGA	<b>A</b> :	120
	TTT	AATT	GGC (	GAGA	ACCA	CA C	GCC	GAAA'	т та	AAAA	AAAA	AAA	A				:	164

(2) INFORMATION FOR SEQ ID NO:23: /

	(i)	(E (C	UENC LE S) TY C) ST O) TO	ENGTH (PE: (RANI	: 52 nucl EDNE	22 ba .eic ESS:	ase p acid unkr	oairs 1	3							
	(ii)	MOI	LECUI	LE TY	PE:	CDNA	7									
-	(vi)	(I	GINA A) OI	RGANI vi	SM: Tus	por		_			e and	d re	spir	ator	y syn	ıdrome
	(ix)	(2	ATURI A) NZ 3) L(	AME/F			519									·•
	(xi)	SEÇ	QUEN	CE DE	ESCR	IPTI	ON: S	SEQ :	ED N	0:23	:					
			CTA Leu												Lys	48
			GCC Ala 20											Tyr	GCC Ala	96
			TCA Ser												ATA lle	144
			TGT Cys									Tyr			TTT Phe	192
			AAC Asn			Ala					Ala				CTT Leu 80	240
			GTT Val							Trp					Ser	288
				Cys					Arg					Pro	GCC Ala	336

CAT His	CAC His	GTA Val 115	GAA Glu	AGT Ser	GCT Ala	GCA Ala	GGT Gly 120	CTC Leu	CAT His	TCA Ser	ATC Ile	TCA Ser 125	GCG Ala	TCT Ser	GGT Gly	384
AAC Asn	CGA Arg 130	GCA Ala	TAC Tyr	GCT Ala	GTG Val	AGA Arg 135	AAG Lys	CCC Pro	GGA Gly	CTA Leu	ACA Thr 140	TCA Ser	GTG Val	AAC Asn	GGC Gly	432
ACT Thr 145	CTA Leu	GTA Val	CCA Pro	GGA Gly	CTT Leu 150	CGG Arg	AGC Ser	CTC Leu	GTG Val	CTG Leu 155	GGC Gly	GGC Gly	AAA Lys	CGA Arg	GCT Ala 160	480
GTT Val	AAA Lys	CGA Arg	GGA Gly	GTG Val 165	GTT Val	AAC Asn	CTC Leu	GTC Val	AAG Lys 170	TAT Tyr	GGC Gly	CGG Arg	TAA			522
[](2)	INF	ORMA!	TION	FOR	SEQ	ID I	NO:24	4:						·		-
		(i) :	(B)	LEI (	NGTH PE: a	: 17: amin	ERIS' 3 am: 0 ac: line	ino id		s				-		
i ab	(:	ii) 1	MOLE	CULE	TYP	E: p:	rote	in								
	(2	xi) :	SEQUI	ENCE	DES	CRIP'	TION	: SE	Q ID	NO:	24:					
Het D 1	Gly	Gly	Leu	Asp 5	Asp	Phe	Сув	Asn	Asp 10	Pro	Ile			15		
∭ i≟Met ij 1	Gly	Gly	Leu	Asp 5	Asp	Phe	Сув	Asn	Asp 10	Pro	Ile			15		
Het Met D 1 J Leu	Gly Val	Gly Leu	Leu Ala 20	Asp 5 Phe	Asp Ser	Phe	Cys	Asn Tyr 25	Asp 10 Thr	Pro	Ile	Met	Ile 30	15 Tyr		
Het Het Leu	Gly Val Lys	Gly Leu Val 35	Leu Ala 20 Ser	Asp 5 Phe Arg	Asp Ser Gly	Phe Ile Arg	Cys Thr Leu 40	Asn Tyr 25 Leu	Asp 10 Thr	Pro Pro Leu	Ile Ile Leu	Met His 45	Ile 30 Ile	15 Tyr Leu	Ala	
Met 1 Leu Leu Phe	Gly Val Lys Leu 50	Gly Leu Val 35 Asn	Leu Ala 20 Ser Cys	Asp 5 Phe Arg Ser	Asp Ser Gly Phe	Phe Ile Arg Thr 55	Cys Thr Leu 40 Phe	Asn Tyr 25 Leu Gly	Asp 10 Thr Gly	Pro Pro Leu Met	Ile Ile Leu Thr 60	Met His 45 Tyr	Ile 30 Ile Val	15 Tyr Leu His	Ala	
Met 1 1 1 Leu  Leu  Phe  Gln 65	Gly Val Lys Leu 50 Ser	Gly Leu Val 35 Asn	Leu Ala 20 Ser Cys	Asp 5 Phe Arg Ser	Asp Ser Gly Phe Val	Phe Ile Arg Thr 55	Cys Thr Leu 40 Phe	Asn Tyr 25 Leu Gly	Asp 10 Thr Gly Tyr	Pro Leu Met Gly 75	Ile Ile Leu Thr 60	Met His 45 Tyr	Ile 30 Ile Val	15 Tyr Leu His	Ala Ile Phe Leu 80	
Met 1 Leu  Leu  Phe  Gln 65  Leu	Gly Val Lys Leu 50 Ser	Gly Leu Val 35 Asn Thr	Leu Ala 20 Ser Cys Asn Val	Asp 5 Phe Arg Ser Arg Tyr 85	Asp Ser Gly Phe Val 70 Ser	Phe Ile Arg Thr 55 Ala Phe	Cys Thr Leu 40 Phe Leu Thr	Asn Tyr 25 Leu Gly Thr	Asp 10 Thr Gly Tyr Leu Ser 90	Pro Leu Met Gly 75	Ile Leu Thr 60 Ala	Met His 45 Tyr Val	Ile 30 Ile Val Val	Tyr Leu His Ala Thr 95	Ala Ile Phe Leu 80	

	Asn	Arg 130		Tyr	Ala	Val	Arg 135	Lys	Pro	Gly	Leu	Thr 140	Ser	Val	Asn	Gly	
	Thr 145	Leu	Val	Pro	Gly	Leu 150	Arg	Ser	Leu	Val	Leu 155	Gly	Gly	Lys	Arg	Ala 160	
	Val	Lys	Arg	Gly	Val 165	Val	Asn	Leu	Val	Lys 170	Tyr	Gly	Arg				
	(2)	INFO	ORMA'	rion	FOR	SEQ	ID I	NO:25	5:								
		(i)	(1 (1 (0	A) LI B) T	ENGTI (PE : [RAN]	1: 30 nuc. DEDNI	37 ba leic ESS:	ISTIC ase p acic unki nown	pair: d	<b>3</b>							
		(ii)	MO	LECU	LE T	YPE:	cDN	A									:
		(vi)	. (2		RGAN: vi:	ISM: rus	por	cine OLAT				e an	d re	spir	ator	y syn	adrome
		(ix	(,	ATUR A) N B) L	AME/												
		(xi	) SE	QUEN	CE D	ESCR	IPTI	ON:	SEQ	ID N	0:25	:					
ė								CAG Gln			Lys					Pro	48
	ATG Met	GGG Gly	AAT Asn	GGC Gly 20	Gln	CCA Pro	GTC Val	AAT Asn	CAA Gln 25	Leu	TGC Cys	CAG Gln	TTG Leu	CTG Leu 30	Gly	GCA Ala	96
	ATG Met	ATA Ile	AAG Lys 35	Ser	CAG Gln	CGC Arg	CAG Gln	CAA Gln 40	Pro	AGG Arg	GGA Gly	GGA Gly	CAG Gln 45	Ala	AAA Lys	AAG 3 Lya	144
			Pro										Glu			ATC o Ile	192

CGG CAC CAC CTC ACC CAG ACT GAA CGC TCC CTC TGC TTG CAA TCG ATC Arg His His Leu Thr Gln Thr Glu Arg Ser Leu Cys Leu Gln Ser Ile

			Phe					GGA							
			AGT Ser 100												
			ATT Ile												
TAA															
(2)	INF	ORMA'	rion	FOR	SEQ	ID I	NO:2	6 :							
		(i) :	SEQUE (A) (B) (D)	LEI TY	NGTH PE: 8	: 120 amino		ino a id		3					
	(:	ii) 1	MOLE	CULE	TYP	E: p	rote	in			·				
[ <del> </del>	(2	ki) :	SEQUE	ENCE	DES	CRIP'	rion	: SE	Q ID	NO:	26:				
I Met II 1	Ala	Gly	Lys	Asn 5	Gln	Ser	Gln	Lys	Lys 10	Lys	Lys	Ser	Thr	Ala 15	Pro
in Met	Gly	Asn	Gly 20	Gln	Pro	Val	Asn	Gln 25	Leu	Cys	Gln	Leu	Leu 30	Gly	Ala
Met	Ile	Lys 35	Ser	Gln	Arg	Gln	Gln 40		Arg	Gly	Gly	Gln 45	Ala	Lys	Lys
Lys	Lys 50	Pro	Glu	Lys	Pro	His 55	Phe	Pro	Leu	Ala	Ala 60	Glu	Asp	Asp	Ile
Arg 65	His	His	Leu	Thr	Gln 70	Thr	Glu	Arg	Ser	Leu 75	Cys	Leu	Gln	Ser	Ile 80
Gln	Thr	Ala	Phe	Asn 85	Gln	Gly	Ala	Gly	Thr 90	Ala	Ser	Leu	Ser	Ser 95	Ser
Gly	Lys	Val	Ser 100	Phe	Gln	Val	Glu	Phe 105	Met	Leu	Pro	Val	Ala 110	His	Thr
Val	Arg	Leu 115	Ile	Arg	Val	Thr	Ser 120		Ser	Ala	Ser	Gln 125	Gly	Ala	Ser

(2)	INFOR	RMATION FOR SEQ ID NO:27:	
	(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 127 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: unknown  (D) TOPOLOGY: unknown	
	(ii)	MOLECULE TYPE: cDNA	
	(vi)	ORIGINAL SOURCE:  (A) ORGANISM: porcine reproductive and respiratory syndromory virus  (C) INDIVIDUAL ISOLATE: Lelystad	ne
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:27:	
TTTC	ACAG	TC AGGTGAATGG CCGCGATTGG CGTGTGGCCT CTGAGTCACC TATTCAATTA	60
GGGG	CGATC	AC ATGGGGGTCA TACTTAATCA GGCAGGAACC ATGTGACCGA AATTAAAAAA	120
AAA	AAAA		127
i= : (2)	INFO	RMATION FOR SEQ ID NO:28:	
	(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 26 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: unknown  (D) TOPOLOGY: linear	·
	(ii)	MOLECULE TYPE: DNA (genomic)	
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:28:	
GGG	GATCC	GG TATTTGGCAA TGTGTC	26
(2)	INFO	RMATION FOR SEQ ID NO:29:	
	(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 28 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: unknown  (D) TOPOLOGY: linear	
	(ii)	MOLECULE TYPE: DNA (genomic)	

		(xi)	SEQUENCE DESCRIPTION: SEQ/ID NO:	29:	
	GGTG	TTTT(	CC ACGAGAACCG CTTAAGGG		28
	(2)	INFO	RMATION FOR SEQ ID NO:30:		
,		(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 21 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: unknown  (D) TOPOLOGY: linear		
		(ii)	MOLECULE TYPE: DNA (genomic)		
,					
( an		(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:	30:	
		GATCC.	AG AGTTTCAGCG G		21
		INFO	RMATION FOR SEQ ID NO:31:		
		(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 25 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: unknown (D) TOPOLOGY: linear		
		(ii)	MOLECULE TYPE: DNA (genomic)		
į sit		(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:	:31:	
	CAG	TTAGT	CCG ACACGGTCTT AAGGG		25
	(2)	INFO	DRMATION FOR SEQ ID NO:32:		•
		(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 22 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: unknown  (D) TOPOLOGY: linear		
		(ii)	MOLECULE TYPE: DNA (genomic)		
	•				
		(xi)	SEQUENCE DESCRIPTION: SEQ ID NO	:32:	
	GGG	САТСО	CTT GTTAAATATG CC		-23

(2)	INFO	RMATION FOR SEQ ID NO:33:	
	(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 19 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: unknown (D) TOPOLOGY: linear	
	(ii)	MOLECULE TYPE: DNA (genomic)	
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:33:	
CTT	ACGCA	CC ACTTAAGGG	19
(2)	INFO	RMATION FOR SEQ ID NO:34:	
	(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 16 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: unknown  (D) TOPOLOGY: linear	·
Jones Bard Br.	(ii)	MOLECULE TYPE: DNA (genomic)	
i 📥	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:34:	
D AAT	GGGGC	TT CTCCGG	16
(2)	INFO	RMATION FOR SEQ ID NO:35:	
	(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 886 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: unknown  (D) TOPOLOGY: unknown	
	(ii)	MOLECULE TYPE: cDNA	
	(vi)	ORIGINAL SOURCE:  (A) ORGANISM: porcine reproductive and respiratory syndrome virus  (B) STRAIN: Iowa  (C) INDIVIDUAL ISOLATE: ISU-12 (VR 2385/VR 2386)	<b>)</b>

## (xi) SEQUENCE DESCRIPTION: SEQ/ID NO:35: ATGGAGTCGT CCTTAGATGA CTTCTGTCAT GATAGCACGG CTCCACAAAA GGTGCTCTTG 60 GCGTTTTCTA TTACCTACAC GCCAGTGATG ATATATGCCC TAAAGGTGAG TCGCGGCCGA 120 CTGCTAGGGC TTCTGCACCT TTTGGTCTTC CTGAATTGTG CTTTCACCTT CGGGTACATG 180 ACATTCGTGC ACTTCAGAG TACAAATAAG GTCGCGCTCA CTATGGGAGC AGTAGTTGCA 240 CTCCTTTGGG GGGTGTACTC AGCCATAGAA ACCTGGAAAT TCATCACCTC CAGATGCCGT 300 TTGTGCTTGC TAGGCCGCAA GTACATTCTG GCCCCTGCCC ACCACGTTGA AAGTGCCGCA 360 GGCTTTCATC CGATTGCGGC AAATGATAAC CACGCATTTG TCGTCCGGCG TCCCGGCTCC 420 ACTACGGTCA ACGGCACATT GGTGCCCGGG TTAAAAAGCC TCGTGTTGGG TGGCAGAAAA 480 GCTGTTAAAC AGGGAGTGGT AAACCTTGTT AAATATGCCA AATAACACCG GCAAGCAGCA 540 I E GAAGAGAAAG AAGGGGGATG GCCAGCCAGT CAATCAGCTG TGCCAGATGC TGGGTAAGAT 600 IT CATCGCTCAC CAAAACCAGT CCAGAGGCAA GGGACCGGGA AAGAAAAATA AGAAGAAAAA 660 CCCGGAGAAG CCCCATTTCC CTCTAGCGAC TGAAGATGAT GTCAGACATC ACTTTACCCC 720 TAGTGAGCGT CAATTGTGTC TGTCGTCAAT CCAGACCGCC TTTAATCAAG GCGCTGGGAC 780 TTGCACCCTG TCAGATTCAG GGAGGATAAG TTACACTGTG GAGTTTAGTT TGCCTACGCA 840 ١Į TCATACTGTG CGCCTGATCC GCGTCACAGC ATCACCCTCA GCATGA 886

# (2) INFORMATION FOR SEQ ID NO:36:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 886 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: unknown
  - (D) TOPOLOGY: unknown
- (ii) MOLECULE TYPE: cDNA
- (vi) ORIGINAL SOURCE:
  - (A) ORGANISM: porcine reproductive and respiratory syndrome virus
  - (B) STRAIN: Iowa
  - (C) INDIVIDUAL ISOLATE: ISU-1894

## ATGGGGTCGT CCTTAGATGA CTTCTGCCAT GATAGTACGG CTCCACAAAA GGTGCTTTTG 60 GCGTTTTCTA TTACCTACAC GCCAGTGATG ATATATGCCC TAAAGGTGAG TCGCGGCCGA 120 CTGCTAGGGC TTCTGCACCT TTTGATCTTC CTGAATTGTG CTTTCACCTT CGGGTACATG 180 ACATTCGTGC ACTTTCAGAG TACAAATAAG GTCGCGCTCA CTATGGGAGC AGTAGTTGCA 240 CTCCTTTGGG GGGTGTACTC AGCCATAGAA ACCTGGAAAT TCATCACCTC CAGATGCCGT 300 TTGTGCTTGC TAGGCCGCAA GTACATTCTG GCCCCTGCCC ACCACGTTGA AAGTGCCGCA 360 GGCTTTCATC CGATTGCGGC AAATGATAAC CACGCATTTG TCGTCCGGCG TCCCGGCTCC 420 ACTACGGTCA ACGCACATT GGTGCCCGGG TTGAAAAGCC TCGTGTTGGG TGGCAGAAAA 480 © GCTGTTAAAC AGGGAGTGGT AAACCTTGTC AAATATGCCA AATAACAACG GCAAGCAGCA 540 D AGAGAGAAAG AAGGGGGATG GCCAGCCAGT CAATCAGCTG TGCCAGATGC TGGGTAAGAT 600 CATCGCTCAG CAAAACCAGT CCAGAGGCAA GGGACCGGGA AAGAAAAACA AGAAGAAAAA 660 CCCGGAGAAG CCCCATTTTC CTCTAGCGAC TGAAGATGAT GTCAGACATC ACTTCACCCC 720 TAGTGAGCGG CAATTGTGTC TGTCGTCAAT CCAGACCGCC TTTAATCAAG GCGCTGGGAC 780 TTGCACCCTG TCAGATTCAG GGAGGATAAG TTACACTGTG GAGTTTAGTT TGCCAACGCA 840

(2) INFORMATION FOR SEQ ID NO:37:

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- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 886 base pairs

TCATACTGTG CGCTTGATCC GCGTCACAGC ATCACCCTCA GCATGA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:36:

- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: unknown
- (D) TOPOLOGY: unknown
- (ii) MOLECULE TYPE: cDNA
- (vi) ORIGINAL SOURCE:
  - (A) ORGANISM: porcine reproductive and respiratory syndrome virus

- (B) STRAIN: Iowa
- (C) INDIVIDUAL ISOLATE: ISU-22 (VR 2429)

(xi) SEQUENCE DESCRIPTION: SEQ/ID NO:37:

			/			
ATGGGGTCGT	CCTTAGATGA	CTTCTGTCAT	GACAGCACGG	CTCCACAAAA	GGTGCTTTTG	60
GCGTTTTCTA	TTACCTACAC	GCCAGTGATG	ATATATGCCC	TGAAGGTGAG	TCGCGGCCGA	120
CTGCTAGGGC	TTCTGCACCT	TTTGATCTTC	CTGAATTGTG	CTTTCACCTT	CGGGTACATG	180
ACATTCGTGC	ACTTTCAGAG	TACAAATAAG	GTCGCACTCA	CTATGGGAGC	AGTAGTTGCA	240
CTCCTTTGGG	GGGTGTACTC	AGCCATAGAA	ACCTGGAAAT	TCATCACCTC	CAGATGCCGT	300
TTGTGCTTGC	TAGGCCGCAA	GTACATTCTG	GCCCTGCCC	ACCACGTTGA	AAGTGCCGCA	360
GGCTTTCATC	CGATTGCGGC	AAATGATAAC	CACGCATTTG	TCGTTCGGCG	TCCCGGCTCC	420
ACTACGGTCA	ACGGCACATT	GGTGCCCGGG	TTGAAAAGCC	TCGTGTTGGG	TGGCAGAAAA	480
GCTGTTAAAC	AGGGAGTGGT	AAACCTTGTC	AAATATGCCA	AATAACAACG	GTAAGCAGCA	540
GAAGAGAAAG	AAGGGGGATG	GCCAGCCAGT	CAATCAGCTG	TGCCAGATGC	TGGGCAAGAT	600
CATCGCTCAG	CAAAATCAGT	CCAGAGGCAA	GGGACCGGGA	AAGAAAAATA	AGAAGAAAA	660
CCCGGAGAAG	CCCCATTTTC	CTCTAGCGAC	TGAAGATGAT	GTCAGACATC	ACTTTACCCC	720
TAGTGAGCGG	CAATTGTGTC	TGTCGTCAAT	CCAGACCGCC	TTTAATCAAG	GCGCTGGGAC	780
TTGCACCCTG	TCAGATTCAG	GGAGGATAAG	TTACACTGTG	GAGTTTAGTT	TGCCTACGCA	840
TCATACTGTG	CGCCTGATCC	GCGTCACAGC	ATCACCCTCA	GCATGA		886

## (2) INFORMATION FOR SEQ ID NO:38:

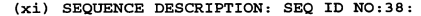
- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 886 base pairs

  - (B) TYPE: nucleic acid (C) STRANDEDNESS: unknown
  - (D) TOPOLOGY: unknown

# (ii) MOLECULE TYPE: cDNA

# (vi) ORIGINAL SOURCE:

- (A) ORGANISM: porcine reproductive and respiratory syndrome virus
- (B) STRAIN: Iowa
- (C) INDIVIDUAL ISOLATE: ISU-79



ATGGGGTCGT	CCTTAGATGA	CTTCTGTTAT	GATAGTACGG	CTCCACAAAA	GGTGCTTTTG	60
GCATTTTCTA	TTACCTACAC	GCCAGTAATG	ATATATGCCC	TAAAGGTGAG	TCGCGGCCGA	120
CTGCTAGGGC	TTCTGCACCT	TTTGATTTTC	CTGAACTGTG	CTTTCACCTT	CGGGTACATG	180
ACATTCATGC	ACTTTCAGAG	TACAAATAAG	GTCGCGCTCA	CTATGGGAGC	AGTAGTTGCA	240
CTCCTTTGGG	GGGTGTACTC	AGCCATAGAA	ACCTGGAAAT	TCATCACCTC	CAGATGCCGT	300
TTGTGCTTGC	TAGGCCGCAA	GTACATTCTG	GCCCCTGCCC	ACCACGTTGA	AAGTGCCGCA	360
GGCTTTCATC	CGATTGCGGC	AAATGATAAC	CACGCATTTG	TCGTCCGGCG	TCCCGGCTCC	420
ACTACGGTCA	ACGGCACATT	GGTGCCCGGG	TTGAAAAGCC	TCGTGTTGGG	TGGCAGAAAA	480
GCTGTTAAAC	AGGGAGTGGT	AAACCTTGTC	AAATATGCCA	AATAACAACG	GCAAGCAGCA	540
GAAGAGAAAG	AAGGGGGATG	GCCAGCCAGT	CAATCAGCTG	TGCCAGATGC	TGGGTAAGAT	600
CATCGCCCAG	CAAAACCAGT	CTAGAGGCAA	GGGACCGGGA	AAGAAAAATA	AGAAGAAAAA	660
CCCGGAGAAG	CCCCATTTTC	CTCTAGCGAC	TGAAGATGAT	GTCAGACATC	ACTTTACCCC	720
TAGTGAGCGG	CAATTGTGTC	TGTCGTCAAT	CCAAACTGCC	TTTAATCAAG	GCGCTGGGAC	780
TTGCACCCTG	TCAGATTCAG	GGAGGATAAG	TTACACTGTG	GAGTTTAGTT	TGCCTACGCA	840
TCATACTGTG	CGCTTGATCC	GCGTCACAGC	ATCACCCTCA	GCATGA		886

# (2) INFORMATION FOR SEQ ID NO:39:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 886 base pairs

  - (B) TYPE: nucleic acid (C) STRANDEDNESS: unknown
  - (D) TOPOLOGY: unknown

# (ii) MOLECULE TYPE: cDNA

## (vi) ORIGINAL SOURCE:

- (A) ORGANISM: porcine reproductive and respiratory syndrome virus
- (B) STRAIN: Iowa
- (C) INDIVIDUAL ISOLATE: ISU-55 (VR 2430)

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:39:

		•				
ATGGGGTCGT	CCTTAGATGA	CTTCTGCCAT	GATAGCACGG	CTCCACAAAA	GGTGCTTTTG	60
GCGTTCTCTA	TTACCTACAC	GCCAGTGATG	ATATATGCCC	TAAAAGTAAG	TCGCGGCCGA	120
CTGCTAGGGC	TTCTGCACCT	TTTGATCTTC	CTAAATTGTG	CTTTCACCTT	CGGGTACATG	180
ACATTCGTGC	ACTTTCAGAG	CACAAACAAG	GTCGCGCTCA	CTATGGGAGC	AGTAGTTGCA	240
CTCCTTTGGG	GGGTGTACTC	AGCCATAGAA	ACCTGGAAAT	TCATCACCTC	CAGATGCCGT	300
TTGTGCTTGC	TAGGCCGCAA	GTACATTTTG	GCCCTGCCC	ACCACGTTGA	AAGTGCCGCA	360
GGCTTTCATC	CGATAGCGGC	AAATGATAAC	CACGCATTTG	TCGTCCGGCG	TCCCGGCTCC	420
	ACGGCACATT	GGTGCCCGGG	TTGAAAAGCC	TCGTGTTGGG	TGGCAGAAAA	480
	AGGGAGTGGT	AAACCTTGTT	AAATATGCCA	AATAACAACG	GCAAGCAGCA	540
GAAGAAAAG	AAGGGGGATG	GCCAGCCAGT	CAATCAGCTG	TGCCAGATGC	TGGGTAAGAT	600
CATCGCTCAG	CAAAACCAGT	CCAGAGGCAA	GGGACCGGGA	AAGAAAAACA	AGAAGAAAAA	660
CCCGGAGAAG	CCCCATTTTC	CTCTAGCGAC	TGAAGATGAT	GTCAGACATC	ACTTCACCTC	720
TGGTGAGCGG	CAATTGTGTC	TGTCGTCAAT	CCAGACAGCC	TTTAATCAAG	GCGCTGGAAC	780
TTGTACCCTG	TCAGATTCAG	GGAGGATAAG	TTACACTGTG	GAGTTTAGTT	TGCCGACGCA	840
TCATACTGTG	CGCTTGATCC	GCGTCACAGC	GTCACCCTCA	GCATGA		886

# (2) INFORMATION FOR SEQ ID NO:40:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 886 base pairs

  - (B) TYPE: nucleic acid (C) STRANDEDNESS: unknown
  - (D) TOPOLOGY: unknown
- (ii) MOLECULE TYPE: cDNA
- (vi) ORIGINAL SOURCE:
  - (A) ORGANISM: porcine reproductive and respiratory syndrome virus
  - (B) STRAIN: Iowa
  - (C) INDIVIDUAL ISOLATE: ISU-3927 (VR 2431)

ATGGGGTCGT CCCTAGACGA CTTTTGCAAT GATAGCACGG CTCCACAAAA GGTGCTTTTG 60 GCGTTTTCTA TTACCTACAC GCCGGTGATG ATATATGCTC TAAAGGTAAG TCGCGGCCGA 120 CTGCTAGGGC TTCTGCACCT TTTGATTTTT CTGAATTGTG CTTTTACTTT CGGGTACATG 180 ACATTCGTGC ACTTTGAGAG CACAAATAGG GTCGCGCTCA CTATGGGAGC AGTAGTCGCA 240 CTTCTCTGGG GGGTGTACTC AGCCATAGAA ACCTGGAAAT TCATCACCTC CAGATGCCGT 300 TTGTGCTTGC TAGGCCGCAA GTACATTCTG GCCCCTGCCC ACCACGTTGA GAGTGCCGCA 360 GGCTTTCATC CGATTGCGGC AAATGATAAC CACGCATTTG TCGTCCGGCG TCCCGGCTCC 420 ACTACGGTTA ACGGCACATT GGTGCCCGGG TTGAGAAGCC TCGTGTTGGG TGGCAAAAAA 480 ١Đ GCTGTTAAGC AGGGAGTGGT AAACCTTGTT AAATATGCCA AATAACAACG GCAAGCAGCA 540 I GAAGAAAAG AAGGGGGATG GCCAGCCAGT CAATCAGCTC TGCCAAATGC TGGGTAAGAT 600 CATCGCCCAG CAAAACCAGT CCAGAGGTAA GGGACCGGGA AAGAAAAATA AGAAGAAAAA 660 CCCGGAGAAG CCCCATTTTC CTCTAGCGAC TGAAGATGAT GTCAGACATC ACTTCACCCC 720 CAGTGAGCGG CAATTGTGTC TGTCGTCAAT CCAGACTGCC TTTAATCAGG GCGCTGGGAC 780 W CTGTATCCTA TCAGATTCAG GGAGGATAAG TTACACTGTG GAGTTTAGTT TGCCGACGCA 840 14 ٠D TCATACTGTG CGCCTGATTC GCGTCACGGC ACCACCCTCA GCATGA 886

- (2) INFORMATION FOR SEQ ID NO:41:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 898 base pairs

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:40:

- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: unknown
- (D) TOPOLOGY: unknown
- (ii) MOLECULE TYPE: cDNA
- (vi) ORIGINAL SOURCE:
  - (A) ORGANISM: porcine reproductive and respiratory syndrome virus
  - (C) INDIVIDUAL ISOLATE: Lelystad
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:41:

ATGGGAGGCC TAGACGATTT TTGCAACGAT CCTATCGCCG CACAAAAGCT CGTGCTAGCC

	TTTAGCATCA	CATACACACC	TATAATGATA	TACGCCCTTA	AGGTGTCACG	CGGCCGACTC	120
	CTGGGGCTGT	TGCACATCCT	AATATTTCTG	AACTGTTCCT	TTACATTCGG	ATACATGACA	180
	TATGTGCATT	TTCAATCCAC	CAACCGTGTC	GCACTTACCC	TGGGGGCTGT	TGTCGCCCTT	240
	CTGTGGGGTG	TTTACAGCTT	CACAGAGTCA	TGGAAGTTTA	TCACTTCCAG	ATGCAGATTG	300
	TGTTGCCTTG	GCCGGCGATA	CATTCTGGCC	CCTGCCCATC	ACGTAGAAAG	TGCTGCAGGT	360
	CTCCATTCAA	TCTCAGCGTC	TGGTAACCGA	GCATACGCTG	TGAGAAAGCC	CGGACTAACA	420
	TCAGTGAACG	GCACTCTAGT	ACCAGGACTT	CGGAGCCTCG	TGCTGGGCGG	CAAACGAGCT	480
	GTTAAACGAG	GAGTGGTTAA	CCTCGTCAAG	TATGGCCGGT	AAAAACCAGA	GCCAGAAGAA	540
150		ACAGCTCCGA	TGGGGAATGG	CCAGCCAGTC	AATCAACTGT	GCCAGTTGCT	600
	GGGTGCAATG	ATAAAGTCCC	AGCGCCAGCA	ACCTAGGGGA	GGACAGGCCA	AAAAGAAAA	660
	GCCTGAGAAG	CCACATTTTC	CCCTGGCTGC	TGAAGATGAC	ATCCGGCACC	ACCTCACCCA	720
		TCCCTCTGCT	TGCAATCGAT	CCAGACGGCT	TTCAATCAAG	GCGCAGGAAC	780
	TGCGTCGCTT	TCATCCAGCG	GGAAGGTCAG	TTTTCAGGTT	GAGTTTATGC	TGCCGGTTGC	840
	TCATACAGTG	CGCCTGATTC	GCGTGACTTC	TACATCCGCC	AGTCAGGGTG	CAAGTTAA	898

(2) INFORMATION FOR SEQ ID NO:42:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 525 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: unknown
- (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: cDNA
- (vi) ORIGINAL SOURCE:
  - (A) ORGANISM: porcine reproductive and respiratory syndrome virus
  - (B) STRAIN: Iowa
  - (C) INDIVIDUAL ISOLATE: ISU-1894
- (ix) FEATURE:
  - (A) NAME/KEY: CDS
  - (B) LOCATION: 1..522

# (xi) SEQUENCE DESCRIPTION: SEQ/ID NO:42:

						•			/									
														GCT Ala		CAA Gln	•	48
														ATG Met 30	Ile	TAT Tyr		96
														CAC His		TTG Leu	1	44
i maj	Ile							Thr						TTC Phe		CAC His	. 1	92
H	TTT Phe	CAG Gln	AGT Ser	ACA Thr	AAT Asn	AAG Lys 70	GTC Val	GCG Ala	CTC Leu	ACT Thr	ATG Met 75	GGA Gly	GCA Ala	GTA Val	GTT Val	GCA Ala 80	2	40
																ACC Thr	2	88
										Arg				CTG Leu 110	Ala	CCT Pro	3	36
	GCC Ala	CAC His	CAC His 115	GTT Val	GAA Glu	AGT Ser	GCC Ala	GCA Ala 120	Gly	TTT Phe	CAT His	CCG Pro	ATT Ile 125	GCG Ala	GCA Ala	AAT Asn	3	84
								Arg					Thr	ACG Thr		AAC Asn	4	32
		Thr					Leu					Leu		GGC Gly		AAA Lys 160	4	80
						Val					Lys			AAA Lys			5	522
	TAA				•												5	525

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# (2) INFORMATION FOR SEQ ID NO:43:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 174 amino acids
  - (B) TYPE: amino acid
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:43:

Met Gly Ser Ser Leu Asp Asp Phe Cys His Asp Ser Thr Ala Pro Gln
1 10 15

Lys Val Leu Leu Ala Phe Ser Ile Thr Tyr Thr Pro Val Met Ile Tyr 20 25 30

Ala Leu Lys Val Ser Arg Gly Arg Leu Leu Gly Leu Leu His Leu Leu 60 45

Ile Phe Leu Asn Cys Ala Phe Thr Phe Gly Tyr Met Thr Phe Val His 50 55 60

Phe Gln Ser Thr Asn Lys Val Ala Leu Thr Met Gly Ala Val Val Ala 65 70 75 80

Leu Leu Trp Gly Val Tyr Ser Ala Ile Glu Thr Trp Lys Phe Ile Thr 85 90 95

Ser Arg Cys Arg Leu Cys Leu Leu Gly Arg Lys Tyr Ile Leu Ala Pro 100 105 110

Ala His His Val Glu Ser Ala Ala Gly Phe His Pro Ile Ala Ala Asn 115 120 125

Asp Asn His Ala Phe Val Val Arg Arg Pro Gly Ser Thr Thr Val Asn 130 135 140

Gly Thr Leu Val Pro Gly Leu Lys Ser Leu Val Leu Gly Gly Arg Lys 145 150 155 160

Ala Val Lys Gln Gly Val Val Asn Leu Val Lys Tyr Ala Lys 165 170

### (2) INFORMATION FOR SEQ ID NO:44:

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- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 525 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: unknown
  - (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA (vi) ORIGINAL SOURCE: (A) ORGANISM: porcine reproductive and respiratory syndrome virus (B) STRAIN: Iowa (C) INDIVIDUAL ISOLATE: ISU-22 (VR 2429) (ix) FEATURE: (A) NAME/KEY: CDS (B) LOCATION: 1..522 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:44: ATG GGG TCG TCC TTA GAT GAC TTC TGT CAT GAC AGC ACG GCT CCA CAA 48 Met Gly Ser Ser Leu Asp Asp Phe Cys His Asp Ser Thr Ala Pro Gln AAG GTG CTT TTG GCG TTT TCT ATT ACC TAC ACG CCA GTG ATG ATA TAT 96 Lys Val Leu Leu Ala Phe Ser Ile Thr Tyr Thr Pro Val Met Ile Tyr 20 25 GCC CTG AAG GTG AGT CGC GGC CGA CTG CTA GGG CTT CTG CAC CTT TTG 144 Ala Leu Lys Val Ser Arg Gly Arg Leu Leu Gly Leu Leu His Leu Leu 35 ATC TTC CTG AAT TGT GCT TTC ACC TTC GGG TAC ATG ACA TTC GTG CAC 192 Ile Phe Leu Asn Cys Ala Phe Thr Phe Gly Tyr Met Thr Phe Val His 50 HTTT CAG AGT ACA AAT AAG GTC GCA CTC ACT ATG GGA GCA GTA GTT GCA 240 Phe Gln Ser Thr Asn Lys Val Ala Leu Thr Met Gly Ala Val Val Ala 65 70 CTC CTT TGG GGG GTG TAC TCA GCC ATA GAA ACC TGG AAA TTC ATC ACC 288 Leu Leu Trp Gly Val Tyr Ser Ala Ile Glu Thr Trp Lys Phe Ile Thr 85 TCC AGA TGC CGT TTG TGC TTG CTA GGC CGC AAG TAC ATT CTG GCC CCT 336 Ser Arg Cys Arg Leu Cys Leu Leu Gly Arg Lys Tyr Ile Leu Ala Pro

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100

115 120 GAT AAC CAC GCA TTT GTC GTT CGG CGT CCC GGC TCC ACT ACG GTC AAC 432 Asp Asn His Ala Phe Val Val Arg Arg Pro Gly Ser Thr Thr Val Asn 130 140 135

384

105

GCC CAC CAC GTT GAA AGT GCC GCA GGC TTT CAT CCG ATT GCG GCA AAT

Ala His His Val Glu Ser Ala Ala Gly Phe His Pro Ile Ala Ala Asn

(									AGC/( Ser							
									CTT ( Leu							
	TAA															
	(2)	INFO	RMAI	NOI	FOR	SEQ	ID N	10:45	5:							
			(i) S	(A)	LEI TYI	NGTH PE: 8		am:			3					
							E: pı									
									: SEÇ							
	1	Gly	Ser	Ser	Leu 5	Asp	Asp	Phe	Cys	His 10	Asp	Ser	Thr	Ala	Pro 15	Gln
	Lys	Val	Leu	Leu 20	Ala	Phe	Ser	Ile	Thr 25	Tyr	Thr	Pro	Val	Met 30	Ile	Tyr
·D		Leu	Lys 35	Val	Ser	Arg	Gly	Arg 40	Leu	Leu	Gly	Leu	Leu 45	His	Leu	Leu
	Ile	Phe 50		Asn	Сув	Ala	Phe 55	Thr	Phe	Gly	Tyr	Met 60	Thr	Phe	Val	His
	Phe 65		Ser	Thr	Asn	Lys 70	Val	Ala	Leu	Thr	Met 75	Gly	Ala	Val	Val	Ala 80
	Leu	Leu	Trp	Gly	Val 85	Tyr	Ser	Ala	Ile	Glu 90	Thr	Trp	Lys	Phe	Ile 95	Thr
	Ser	Arg	Сув	Arg 100	Leu	Cys	Leu	Leu	Gly 105		Lys	Tyr	Ile	Leu 110	Ala	Pro
-	Ala	His	His 115		Glu	Ser	Ala	Ala 120		Phe	His	Pro	Ile 125		Ala	Asn
	Asp	Asn 130		Ala	Phe	Val	Val 135		Arg	Pro	Gly	Ser 140	Thr	Thr	Val	Asn
	Gly 145		Leu	Val	Pro	Gly 150		Lys	Ser	Leu	Val 155	Leu	Gly	Gly	Arg	Lys 160

Ala Val Lys Gln Gly Val Val Asn Leu Val Lys Tyr Ala Lys

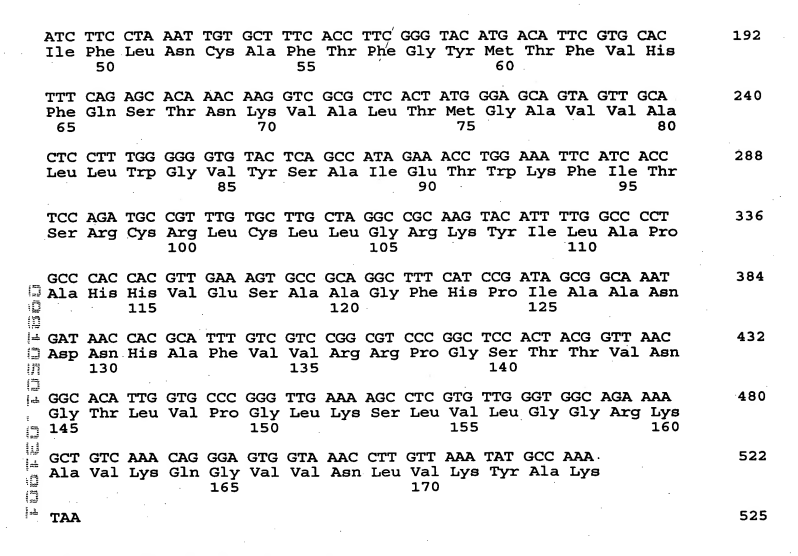
(2) INFORMATION FOR SEQ ID NO:46:

(i) SEQUENCE CHARACTERISTICS:

Leu Leu Trp Gly Val Tyr Ser Ala Ile Glu Thr Trp Lys Phe Ile Thr

		AGA Arg															336
		CAC His		Val													384
		AAC Asn 130														AAC Asn	432
		ACA Thr										Leu				AAA Lys 160	480
		GTT Val									Lys						522
	TAA	INF	ORMA'	rto <b>n</b>	FOR	SEO	TD.	NO : 4	7:							,	525
	(2)	T111 (	J14-11.	1 1 021	1 010				•								
F 1			(i) :	(A (B	) LEI ) TY:	NGTH PE:	: 17 amin	ERIS' 4 am o ac line	ino : id		s						
Cal Gar to the the				(A (B (D	) LEI ) TY: ) TO:	NGTH PE: POLO	: 17 amin GY:	4 am o ac	ino id ar		s						
. In the line in the line in the		. (:	ii) 1	(A (B (D MOLE	) LEI ) TY: ) TO: CULE	NGTH PE: POLO	: 17 amin GY: E: p	4 am o ac line	ino id ar in	acid		47:					
The Control of the State of the	Met 1	(: Gly	ii) l xi) :	(A (B (D MOLE SEQU	) LEI ) TY ) TO  CULE	NGTH PE: POLO  TYP  DES  Asp	: 17 amin GY: E: p CRIP	4 am o ac line rote TION	ino id ar in : SE	acid Q ID	NO:		Thr	Ala	. Pro 15	Gln	
The first the first than the first	1	() ()	ii) ! xi) : Ser	(A (B (D MOLE SEQU Ser	) LEI ) TYI ) TOI CULE ENCE Leu 5	NGTH PE: POLO TYP DES Asp	: 17 amin GY: E: p CRIP Asp	4 am o ac line rote TION	ino id ar in : SE	Q ID Tyr 10	NO:	Ser			15 Ile		
Hart Marie Str. Marie Hart	1 Lys	(; Gly Val	ii)   xi)   Ser Leu	(A (B (D MOLE SEQU Ser Leu 20	) LEI ) TY ) TO CULE ENCE Leu 5	NGTH PE: POLO  TYP  DES Asp	: 17 amin GY: E: p CRIP Asp	4 am o ac line rote TION Phe	ino id ar in : SE Cys Thr 25	Q ID Tyr 10 Tyr	NO: Asp	Ser	Val	Met 30	15		
The last the	1 Lys Ala	() Gly Val Leu	ii)   xi)   Ser Leu Lys 35 Leu	(A (B (D MOLE SEQU Ser Leu 20 Val	) LEI ) TY ) TO CULE ENCE Leu 5 Ala	NGTH PE: POLO TYP DES Asp Phe Arg	: 17 amin GY: E: p CRIP Asp Ser	4 am o ac line rote TION Phe Ile Arg	ino id ar in : SE Cys Thr 25	Q ID Tyr 10 Tyr Leu	NO: Asp Thr	Pro	Val Leu 45	Met 30 His	15 Ile	. Tyr	
	l Lys Ala Ile	(; Gly Val Leu Phe 50	ii) ! xi) : Ser Leu Lys 35 Leu	(A (B (D MOLE SEQU Ser Leu 20 Val	) LEI ) TY ) TO CULE ENCE Leu 5 Ala Ser	NGTH PE: POLO TYP DES Asp Phe Arg	: 17 amin GY: E: p CRIP Asp Ser Gly Phe 55	4 am o ac line rote TION Phe Ile 40	ino id ar in : SE Cys Thr 25 Leu Phe	Q ID Tyr 10 Tyr Leu Gly	NO: Asp Thr Gly	Pro Leu Met	Val Leu 45	Met 30 His	15 Ile	Tyr Leu	

	Ser	Arg	Cys	Arg 100	Leu	Cys	Leu	Leu	Gly 105	Arg	Lys	Tyr	Ile	Leu 110	Ala	Pro	
	Ala	His	His 115	Val	Glu	Ser	Ala	Ala 120	Gly	Phe	His	Pro	Ile 125	Ala	Ala	Asn	
	Asp	Asn 130	His	Ala	Phe	Val	Val 135	Arg	Arg	Pro	Gly	Ser 140	Thr	Thr	Val	Asn	
	Gly 145	Thr	Leu	Val	Pro	Gly 150	Leu	Lys	Ser	Leu	Val 155	Leu	Gly	Gly	Arg	Lys 160	
	Ala	Val	Lys	Gln	Gly 165	Val	Val	Asn	Leu	Val 170	_	Tyr	Ala	Lys			
	(2)	INFO	ORMA!	TION	FOR	SEQ	ID 1	NO:48	B:						٠		
LI THE PERMIT	,	(i)	() ()	A) L: B) T C) S'	CE CI ENGTI YPE: TRANI OPOLO	H: 52 nuci DEDNI	25 ba leic ESS:	ase j acio unki	pair: d	s							
		(ii)	) MO	LECU	LE T	YPE:	cDN	A									
		(vi)	(; (1	A) 0: B) S	vi: TRAI	ISM: rus N: I	poro owa	:	_	rodu SU-5				spir	ator	y syn	drome
-		(ix	(2		E: AME/I OCAT			522			٠.						·
		(xi	) SE	QUEN	CE D	ESCR	IPTI	ON:	SEQ	ID N	0:48	:					
										CAT His 10	Asp					Gln	4
					Ala					_						TAT Tyr	9
				Val					Leu	CTA Leu				His		TTG Leu	14

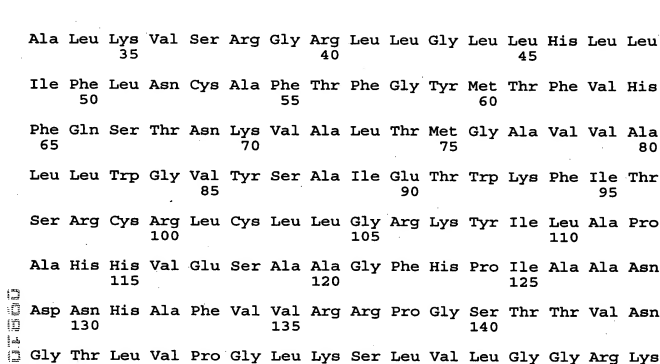


### (2) INFORMATION FOR SEQ ID NO:49:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 174 amino acids
  - (B) TYPE: amino acid
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:49:

Met Gly Ser Ser Leu Asp Asp Phe Cys His Asp Ser Thr Ala Pro Gln
1 5 10 15

Lys Val Leu Leu Ala Phe Ser Ile Thr Tyr Thr Pro Val Met Ile Tyr
20 25 30



Ala Val Lys Gln Gly Val Val Asn Leu Val Lys Tyr Ala Lys
165 170

# (2) INFORMATION FOR SEQ ID NO:50:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 525 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: unknown
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: cDNA
- (vi) ORIGINAL SOURCE:
  - (A) ORGANISM: porcine reproductive and respiratory syndrome virus

155

- (B) STRAIN: Iowa
- (C) INDIVIDUAL ISOLATE: ISU-3927 (VR 2431)
- (ix) FEATURE:

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- (A) NAME/KEY: CDS
- (B) LOCATION: 1..522

# (xi) SEQUENCE DESCRIPTION: SEQ ID NO:50: ATG GGG TCG TCC CTA GAC GAC TTT TGC AAT GAT AGC ACG GCT CCA CAA Met Gly Ser Ser Leu Asp Asp Phe Cys Asn Asp Ser Thr Ala Pro Gln

AAG GTG CTT TTG GCG TTT TCT ATT ACC TAC ACG CCG GTG ATG ATA TAT 96
Lys Val Leu Leu Ala Phe Ser Ile Thr Tyr Thr Pro Val Met Ile Tyr
20 25 30

GCT CTA AAG GTA AGT CGC GGC CGA CTG CTA GGG CTT CTG CAC CTT TTG

Ala Leu Lys Val Ser Arg Gly Arg Leu Leu Gly Leu Leu His Leu Leu

40

45

ATT TTT CTG AAT TGT GCT TTT ACT TTC GGG TAC ATG ACA TTC GTG CAC

Ile Phe Leu Asn Cys Ala Phe Thr Phe Gly Tyr Met Thr Phe Val His

50 55 60

TTT GAG AGC ACA AAT AGG GTC GCG CTC ACT ATG GGA GCA GTA GTC GCA

Phe Glu Ser Thr Asn Arg Val Ala Leu Thr Met Gly Ala Val Val Ala

65 70 75 80

CTT CTC TGG GGG GTG TAC TCA GCC ATA GAA ACC TGG AAA TTC ATC ACC 288
Leu Leu Trp Gly Val Tyr Ser Ala Ile Glu Thr Trp Lys Phe Ile Thr
85 90 95

TCC AGA TGC CGT TTG TGC TTG CTA GGC CGC AAG TAC ATT CTG GCC CCT Ser Arg Cys Arg Leu Cys Leu Leu Gly Arg Lys Tyr Ile Leu Ala Pro 100 105 110

GCC CAC CAC GTT GAG AGT GCC GCA GGC TTT CAT CCG ATT GCG GCA AAT 384
Ala His His Val Glu Ser Ala Ala Gly Phe His Pro Ile Ala Ala Asn
115 120 125

GAT AAC CAC GCA TTT GTC GTC CGG CGT CCC GGC TCC ACT ACG GTT AAC
Asp Asn His Ala Phe Val Val Arg Arg Pro Gly Ser Thr Thr Val Asn
130 135 140

GGC ACA TTG GTG CCC GGG TTG AGA AGC CTC GTG TTG GGT GGC AAA AAA 480 Gly Thr Leu Val Pro Gly Leu Arg Ser Leu Val Leu Gly Gly Lys Lys 150 155 160

GCT GTT AAG CAG GGA GTG GTA AAC CTT GTT AAA TAT GCC AAA Ala Val Lys Gln Gly Val Val Asn Leu Val Lys Tyr Ala Lys 165 170

TAA 525

522

48

192

240

336



- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 174 amino acids
  - (B) TYPE: amino acid
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:51:

Met Gly Ser Ser Leu Asp Asp Phe Cys Asn Asp Ser Thr Ala Pro Gln
1 5 10 15

Lys Val Leu Leu Ala Phe Ser Ile Thr Tyr Thr Pro Val Met Ile Tyr 20 25 30

Ala Leu Lys Val Ser Arg Gly Arg Leu Leu Gly Leu Leu His Leu Leu
35 40 45

File Phe Leu Asn Cys Ala Phe Thr Phe Gly Tyr Met Thr Phe Val His 50 55 60

Phe Glu Ser Thr Asn Arg Val Ala Leu Thr Met Gly Ala Val Val Ala 5 70 75 80

Leu Leu Trp Gly Val Tyr Ser Ala Ile Glu Thr Trp Lys Phe Ile Thr

Ser Arg Cys Arg Leu Cys Leu Leu Gly Arg Lys Tyr Ile Leu Ala Pro

Ala His His Val Glu Ser Ala Ala Gly Phe His Pro Ile Ala Ala Asn 115 120 125

Asp Asn His Ala Phe Val Val Arg Arg Pro Gly Ser Thr Thr Val Asn 130 135 140

Gly Thr Leu Val Pro Gly Leu Arg Ser Leu Val Leu Gly Gly Lys Lys 145 150 155 160

Ala Val Lys Gln Gly Val Val Asn Leu Val Lys Tyr Ala Lys
165 170

- (2) INFORMATION FOR SEQ ID NO:52:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 372 base pairs
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: unknown
    - (D) TOPOLOGY: linear



# (ii) MOLECULE TYPE: cDNA

## (vi) ORIGINAL SOURCE:

- (A) ORGANISM: porcine reproductive and respiratory syndrome virus
- (B) STRAIN: Iowa
- (C) INDIVIDUAL ISOLATE: ISU-1894

# (ix) FEATURE:

(A) NAME/KEY: CDS

(B) LOCATION: 1..369

# (xi) SEQUENCE DESCRIPTION: SEQ ID NO:52:

17												
ATG Met											GGC Gly	48
CAG Gln	 									Ala		96
CAA Gln									Lys			144
AAC Asn								Asp			AGA Arg	192
							Leu		TCA Ser		CAG Gln 80	240
						Thr			GAT Asp		Gly	288
		Thr			Leu				CAT His	Thr	GTG Val	336
	 	 	 GCA Ala	 Pro								372

- (2) INFORMATION FOR SEQ ID NO:53:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 123 amino acids
    - (B) TYPE: amino acid
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: protein
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:53:

Met Pro Asn Asn Gly Lys Gln Gln Lys Arg Lys Gly Asp Gly 1 5 10 15

Gln Pro Val Asn Gln Leu Cys Gln Met Leu Gly Lys Ile Ile Ala Gln 20 25 30

Gln Asn Gln Ser Arg Gly Lys Gly Pro Gly Lys Lys Asn Lys Lys Lys 35 40 45

Asn Pro Glu Lys Pro His Phe Pro Leu Ala Thr Glu Asp Asp Val Arg
50 55 60

His His Phe Thr Pro Ser Glu Arg Gln Leu Cys Leu Ser Ser Ile Gln
65 70 75 80

Thr Ala Phe Asn Gln Gly Ala Gly Thr Cys Thr Leu Ser Asp Ser Gly
85 90 95

Arg Ile Ser Tyr Thr Val Glu Phe Ser Leu Pro Thr His His Thr Val

Arg Leu Ile Arg Val Thr Ala Ser Pro Ser Ala 115 120

- (2) INFORMATION FOR SEQ ID NO:54:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 372 base pairs
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: unknown
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: cDNA
  - (vi) ORIGINAL SOURCE:
    - (A) ORGANISM: porcine reproductive and respiratory syndrome virus
    - (B) STRAIN: Iowa
    - (C) INDIVIDUAL ISOLATE: ISU-22 (VR 2429)

	(ix)	(2		AME/1	KEY:		369	/									
	(xi)	SEC	QUEN	CE DI	ESCR:	IPTIC	ON:	SEQ	ID N	0:54	<b>:</b> .						
					GGT Gly										Gly		48
					CTG Leu									Ala	CAG Gln		96
					GGC Gly										AAA Lys		144
AAC Asn	CCG Pro 50	Glu	AAG Lys	CCC Pro	CAT His	TTT Phe 55	CCT Pro	CTA Leu	GCG Ala	ACT Thr	GAA Glu 60	GAT Asp	GAT Asp	GTC Val	AGA Arg		192
					AGT Ser 70										CAG Gln 80		240
					GGC Gly					Thr					Gly		288
					GTG Val									Thr	GTG Val		336
					ACA Thr												372
(2)	INF	ORMA'	TION	FOR	SEQ	ID :	NO:5	5:								• .	
		(i)	(A (B	) LE ) TY	CHA NGTH PE: POLO	: 12 amin	3 am o ac	ino id		s							

(ii) MOLECULE TYPE: protein



(xi) SEQUENCE DESCRIPTION: SEQ ID NO:55:

Met Pro Asn Asn Asn Gly Lys Gln Gln Lys Arg Lys Lys Gly Asp Gly 15

Gln Pro Val Asn Gln Leu Cys Gln Met Leu Gly Lys Ile Ile Ala Gln 30

Gln Asn Gln Ser Arg Gly Lys Gly Pro Gly Lys Lys Asn Lys Lys Lys Asn Pro Glu Lys Pro His Phe Pro Leu Ala Thr Glu Asp Asp Val Arg 60

His His Phe Thr Pro Ser Glu Arg Gln Leu Cys Leu Ser Ser Ile Gln 65

Thr Ala Phe Asn Gln Gly Ala Gly Thr Cys Thr Leu Ser Asp Ser Gly 90

Arg Ile Ser Tyr Thr Val Glu Phe Ser Leu Pro Thr His His Thr Val 100

Arg Leu Ile Arg Val Thr Ala Ser Pro Ser Ala

- (2) INFORMATION FOR SEQ ID NO:56:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 372 base pairs
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: unknown
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: cDNA
  - (vi) ORIGINAL SOURCE:
    - (A) ORGANISM: porcine reproductive and respiratory syndrome virus
    - (B) STRAIN: Iowa
    - (C) INDIVIDUAL ISOLATE: ISU-79
  - (ix) FEATURE:

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- (A) NAME/KEY: CDS
- (B) LOCATION: 1..369

	(xi)	SEÇ	QUENC	CE DE	ESCR:	IPTI	ON:	SEQ	ID N	0:56	:				
	CCA Pro													GGC Gly	48
	CCA Pro													CAG Gln	96
	AAC Asn													AAA Lys	144
	CCG Pro 50											Asp		AGA Arg	192
CAT THIS - 65															240
ACT														Gly	288
AGG Arg									Leu				Thr	GTG Val	336
	TTG Leu												• •		372

### (2) INFORMATION FOR SEQ ID NO:57:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 123 amino acids
  - (B) TYPE: amino acid
  - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: protein
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:57:

Met Pro Asn Asn Gly Lys Gln Gln Lys Arg Lys Lys Gly Asp Gly
1 5 10 15

Gln Pro Val Asn Gln Leu Cys Gln Met Leu Gly Lys Ile Ile Ala Gln 20 25 30

Gln	Asn	Gln 35	Ser	Arg	Gly	Lys	Gly 40	Pro	Gly	Lys	Lys	Asn 45	Lys	Lys	Lys		
Asn	Pro 50	Glu	Lys	Pro	His	Phe 55	Pro	Leu	Ala	Thr	Glu 60	Asp	Asp	Val	Arg		
His 65	His	Phe	Thr	Pro	Ser 70	Glu	Arg	Gln	Leu	Cys 75	Leu	Ser	Ser	Ile	Gln 80		
Thr	Ala	Phe	Asn	Gln 85	Gly	Ala	Gly	Thr	Cys 90	Thr	Leu	Ser	Asp	Ser 95	Gly		
Arg	Ile	Ser	Tyr 100	Thr	Val	Glu	Phe	Ser 105	Leu	Pro	Thr	His	His 110	Thr	Val		
	Leu	Ile 115	Arg	Val	Thr	Ala	Ser 120	Pro	Ser	Ala							
[] [j(2)	INFO	ORMA!	rion	FOR	SEQ	ID I	NO:58	8:									
	(1)	() ()	A) LI B) T C) S'	ENGTI YPE : IRANI	HARAGE TO THE TOTAL TO THE TOTAL TOT	72 ba leic ESS:	ase j acio unki	pair: d	<b>.</b>								
1	.(ii)	MO:	LECU	LE T	YPE:	CDN	A.										
	(vi)	()	A) 01 B) S'	RGAN vi: TRAI	rus N: I	por owa					e and	:	spir	ator	y syn	drome	:
	(ix	(.		AME/	KEY: ION:		369						·				
	(xi	) SE	QUEN	CE D	ESCR	IPTI	ON:	SEQ	ID N	0:58	:					,	
										Lys					Gly		4.8
	CCA Pro														CAG Gln		96

														AAG i Lys		144
														GTC Z Val		192
														ATC (		240
ACA Thr	GCC Ala	TTT Phe	AAT Asn	CAA Gln 85	GGC Gly	GCT Ala	GGA Gly	ACT Thr	TGT Cys 90	ACC Thr	CTG Leu	TCA Ser	GAT Asp	TCA Ser 95	GGG Gly	288
AGG DArg	ATA Ile	AGT Ser	TAC Tyr 100	ACT Thr	GTG Val	GAG Glu	TTT Phe	AGT Ser 105	TTG Leu	CCG Pro	ACG Thr	CAT His	CAT His 110		GTG Val	336
=CGC																372
[4 , (2)			SEQUI	ENCE LEI	SEQ CHAI NGTH PE: 8	RACT	ERIS	TICS ino		<b>s</b>						
F. L. C.			(D)	ŢOI	POLO	GY:	line	ar								
					TYPI	_			Q ID	NO:	59:					
Met 1	Pro	Asn	Asn	Asn 5	Gly	Lys	Gln	Gln	Lys 10		Lys	Lys	Gly	Asp 15	Gly	
Gln	Pro	Val	Asn 20	Gln	Leu	Cys	Gln	Met 25		Gly	Lys	Ile	Ile 30	Ala	Gln	
Gln	Asn	Gln 35	Ser	Arg	Gly	Lys	Gly 40	Pro	Gly	Lys	Lys	Asn 45		Lys	Lys	
Asn	Pro 50	Glu	Lys	Pro	His	Phe 55	Pro	Leu	Ala	Thr	Glu 60	_	Asp	Val	Arg	
His 65	His	Phe	Thr	Ser	Gly 70	Glu	Arg	Gln	Leu	Cys 75		Ser	Ser	Ile	Gln 80	

Thr Ala Phe Asn Gln Gly Ala Gly Thr Cys Thr Leu Ser Asp Ser Gly 90 Arg Ile Ser Tyr Thr Val Glu Phe Ser Leu Pro Thr His His Thr Val 100 105 Arg Leu Ile Arg Val Thr Ala Ser Pro Ser Ala (2) INFORMATION FOR SEQ ID NO:60: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 372 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: unknown (D) TOPOLOGY: linear (ii) MOLECULE TYPE: cDNA ٠Ō Ħ (vi) ORIGINAL SOURCE: (A) ORGANISM: porcine reproductive and respiratory syndrome lЛ virus ij (B) STRAIN: Iowa (C) INDIVIDUAL ISOLATE: ISU-3927 (VR 2431) į÷, IJ (ix) FEATURE: Ш (A) NAME/KEY: CDS 1 (B) LOCATION: 1..369 : C (xi) SEQUENCE DESCRIPTION: SEQ ID NO:60: 48 Met Pro Asn Asn Asn Gly Lys Gln Gln Lys Lys Lys Gly Asp Gly 1 CAG CCA GTC AAT CAG CTC TGC CAA ATG CTG GGT AAG ATC ATC GCC CAG .96 Gln Pro Val Asn Gln Leu Cys Gln Met Leu Gly Lys Ile Ile Ala Gln CAA AAC CAG TCC AGA GGT AAG GGA CCG GGA AAG AAA AAT AAG AAG AAA 144 Gln Asn Gln Ser Arg Gly Lys Gly Pro Gly Lys Lys Asn Lys Lys Lys 35 AAC CCG GAG AAG CCC CAT TTT CCT CTA GCG ACT GAA GAT GAT GTC AGA 192 Asn Pro Glu Lys Pro His Phe Pro Leu Ala Thr Glu Asp Asp Val Arg

60

55

50

	CAC His															240
ACT Thr	GCC Ala	TTT Phe	AAT Asn	CAG Gln 85	GGC Gly	GCT Ala	GGG Z	ACC Thr	TGT Z Cys 90	ATC ( Ile	CTA : Leu	rca G Ser	Asp	CA G Ser 95	GG Gly	288
	ATA Ile															. 336
	CTG Leu										TGA					372
(2)	INFO	RMA	rion	FOR	SEQ	ID 1	10:61	Ĺ:								
		(i) s	(A)	LEI TY	CHAINGTH PE: 6	: 123	am: ac:	ino i id		3				-		
i.d i.e.	(:	Li) !	MOLE	CULE	TYPI	E: p	rote:	in								,
: ; ===																
	- (3	Ki) S	SEQU	ENCE	DES	CRIP'	rion	: SE	Q ID	NO:	61:					
Met												Lys	Gly	Asp 15	Gly	
Met		Asn	Asn	Asn 5	Gly	Lys	Gln	Gln	Lys 10	Lys	Lys			12		
Met 1 Eln	Pro	Asn Val	Asn 20	Asn 5 Gln	Gly	Cys	Gln	Gln Met 25	Lys 10 Leu	Lys	Lys	Ile	Ile 30	Ala	Gln	
Met 1 Cln	Pro	Asn Val Gln 35	Asn 20 Ser	Asn 5 Gln Arg	Gly Leu Gly	Lys Cys Lys	Gln Gln Gly 40	Gln Met 25 Pro	Lys 10 Leu Gly	Lys Gly Lys	Lys	Ile Asn 45 Asp	Ile 30 Lys	Ala Lys	Gln Lys	
Met I Gln Gln Asn	Pro Pro Asn Pro 50 His	Asn Val Gln 35 Glu	Asn 20 Ser Lys	Asn 5 Gln Arg	Gly Leu Gly His	Lys Cys Lys Phe 55	Gln Gln Gly 40 Pro	Met 25 Pro	Lys 10 Leu Gly	Lys Gly Lys Thr	Lys Lys Glu 60 Leu	Asn 45 Asp	Ile 30 Lys Asp	Ala Lys Val	Gln Lys Arg	
Met I Gln Asn His	Pro Pro Asn Pro 50 His	Asn Val Gln 35 Glu Phe	Asn 20 Ser Lys	Asn 5 Gln Arg Pro	Gly Leu Gly His Ser 70 Gly	Lys Cys Lys Phe 55 Glu	Gln Gly 40 Pro	Gln Met 25 Pro Leu Gln	Lys 10 Leu Gly Ala Leu	Lys Gly Lys Thr Cys 75	Lys Lys Glu 60 Leu	Ile Asn 45 Asp	Ile 30 Lys Asp	Ala Lys Val	Gln Lys Arg Gln 80 Gly	
Met I Gln Asn His 65	Pro Pro Asn Pro 50	Asn Val Gln 35 Glu Phe	Asn 20 Ser Lys Thr	Asn 5 Gln Arg Pro Pro Gln 85	Gly Leu Gly His Ser 70 Gly	Lys Cys Lys Phe 55 Glu Ala	Gln Gly 40 Pro Arg	Gln Met 25 Pro Leu Gln	Lys 10 Leu Gly Ala Leu Cys 90	Lys Gly Lys Thr Cys 75	Lys Lys Glu 60 Leu	Ile Asn 45 Asp Ser	Ile 30 Lys Asp Ser	Ala Lys Val Ile Ser 95 Thr	Gln Lys Arg Gln 80 Gly	

(2) INFORMATION FOR SEQ ID NO:62: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 7 amino acids (B) TYPE: amino acid (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (xi) SEQUENCE DESCRIPTION: SEQ ID NO:62: Lys Lys Ser Thr Ala Pro Met INFORMATION FOR SEQ ID NO:63: :0 (i) SEQUENCE CHARACTERISTICS: I (A) LENGTH: 4 amino acids (B) TYPE: amino acid (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (xi) SEQUENCE DESCRIPTION: SEQ ID NO:63: Ala Ser Gln Gly INFORMATION FOR SEQ ID NO:64: (2) (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 240 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: Other nucleic acid; (A) DESCRIPTION: DNA (synthetic) (vi) ORIGINAL SOURCE: (A) ORGANISM: porcine reproductive and respiratory syndrome virus (B) STRAIN: Iowa (C) INDIVIDUAL ISOLATE: ISU-12 (VR 2385/VR 2386)

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:64:

TCTTCTTGCC TTTTCTATGC TTCTGAGATG AGTGAAAAGG GATTTAAGGT GGTATTTGGC 60 120 AATGTGTCAG GCATCGTGGC AGTGTGCGTC AACTTCACCA GTTACGTCCA ACATGTCAAG GAATTTACCC AACGTTCCTT GGTAGTTGAC CATGTGCGGC TGCTCCATTT CATGACGCCC 180 GAGACCATGA GGTGGGCAAC TGTTTTAGCC TGTCTTTTTA CCATTCTGTT GGCAATTTGA 240 (2) INFORMATION FOR SEQ ID NO:65: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 1799 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: unknown (D) TOPOLOGY: unknown Ď (ii) MOLECULE TYPE: cDNA (vi) ORIGINAL SOURCE: (A) ORGANISM: porcine reproductive and respiratory syndrome virus (B) STRAIN: Iowa (C) INDIVIDUAL ISOLATE: ISU-12 (VR 2385/VR 2386) (xi) SEQUENCE DESCRIPTION: SEQ ID NO:65: ٠Ō CCTGAATTGA GATGAAATGG GGTCTATGCA AAGCCTTTTT GACAAAATTG GCCAACTTTT 60 TGTGGATGCT TTCACGGAGT TCTTGGTGTC CATTGTTGAT ATCATTATAT TTTTGGCCAT 120 TTTGTTTGGC TTCACCATCG CAGGTTGGCT GGTGGTCTTT TGCATCAGAT TGGTTTGCTC 180 240 CGCGATACTC CGTGCGCGCC CTGCCATTCA CTCTGAGCAA TTACAGAAGA TCCTATGAGG CCTTTCTCT TCAGTGCCAG GTGGACATTC CCACCTGGGG AACTAAACAT CCTTTGGGGA 300 TGCTTTGGCA CCATAAGGTG TCAACCCTGA TTGATGAAAT GGTGTCGCGT CGAATGTACC 360 GCATCATGGA AAAAGCAGGA CAGGCTGCCT GGAAACAGGT AGTGAGCGAG GCTACGCTGT 420 CTCGCATTAG TAGTTTGGAT GTGGTGGCTC ATTTTCAGCA TCTTGCCGCC ATTGAAGCCG 480

AGACCTGTAA ATATCTGGCC TCTCGGCTGC CCATGCTACA CCACCTGCGC ATGACAGGGT

CAAATGTAAC CATAGTGTAT AATAGTACTT TGAATCAGGT GTTTGCTGTT TTCCCAACCC

CTGGTTCCCG GCCAAAGCTT CATGATTTCC AGCAATGGCT AATAGCTGTA CATTCCTCTA

540

600

660



### (2) INFORMATION FOR SEQ ID NO:66:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 771 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: unknown
  - (D) TOPOLOGY: linear

#### (ii) MOLECULE TYPE: cDNA

- (vi) ORIGINAL SOURCE:
  - (A) ORGANISM: porcine reproductive and respiratory syndrome virus
  - (B) STRAIN: Iowa



## (C) INDIVIDUAL ISOLATE: ISU-12 (VR 2385/VR 2386)

## (ix) FEATURE:

(A) NAME/KEY: CDS
(B) LOCATION: 1..768

### (xi) SEQUENCE DESCRIPTION: SEQ ID NO:66:

	(XI)	SEÇ	OFNC	JE DE	iscr.	LPTI	JN: 1	SEQ .	א מז	J:66	•					
														AAC 7 Asn 15		48
														TCA '		96
TAT								Ser					Gly	TGG '		144
												Val		GCC Ala		192
CCA Pro	TTC Phe	ACT Thr	CTG Leu	AGC Ser	AAT Asn 70	TAC Tyr	'AGA Arg	AGA Arg	TCC Ser	TAT Tyr 75	Glu	GCC Ala	TTT Phe	CTC Leu	TCT Ser 80	240
										Thr				TTG Leu 95		288
				His					Leu					GTG : Val		336
			Tyr					ı Lys					a Ala	TGG Trp		384
		Val					Let					Se		GAT 1 Asp		432
	Ala					Let					ı Ala			TGT c Cys	AAA Lys 160	480

													ACA Thr 175		528
													TTT Phe		576
								Pro				Phe	CAG Gln	CAA Gln	624
											Val		GCT Ala	TCT Ser	672
TGT Cys 225										Pro			CGT Arg		720
					Trp				Phe				TCA Ser 255	Arg	768
TGA															771
[](2)	INF	ORMA	TION	FOR	SEQ	ID	<b>NO:</b> 6	7:							
		(i)	(B	) LE ) TY		: 25 amin	6 am	ino	ls				•		

- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:67:

(D) TOPOLOGY: linear

Met Lys Trp Gly Leu Cys Lys Ala Phe Leu Thr Lys Leu Ala Asn Phe 1 5 10 15

Leu Trp Met Leu Ser Arg Ser Ser Trp Cys Pro Leu Leu Ile Ser Leu 20 25 30

Tyr Phe Trp Pro Phe Cys Leu Ala Ser Pro Ser Gln Val Gly Trp Trp 35 40 45

Ser Phe Ala Ser Asp Trp Phe Ala Pro Arg Tyr Ser Val Arg Ala Leu 50 60

Pro Phe Thr Leu Ser Asn Tyr Arg Arg Ser Tyr Glu Ala Phe Leu Ser 65 70 75 80

Gln Cys Gln Val Asp Ile Pro Thr Trp Gly Thr Lys His Pro Leu Gly Met Leu Trp His His Lys Val Ser Thr Leu Ile Asp Glu Met Val Ser Arg Arg Met Tyr Arg Ile Met Glu Lys Ala Gly Gln Ala Ala Trp Lys 120 Gln Val Val Ser Glu Ala Thr Leu Ser Arg Ile Ser Ser Leu Asp Val Val Ala His Phe Gln His Leu Ala Ala Ile Glu Ala Glu Thr Cys Lys 160 150 155 Tyr Leu Ala Ser Arg Leu Pro Met Leu His His Leu Arg Met Thr Gly 170 Ser Asn Val Thr Ile Val Tyr Asn Ser Thr Leu Asn Gln Val Phe Ala 180 185 Val Phe Pro Thr Pro Gly Ser Arg Pro Lys Leu His Asp Phe Gln Gln 195 Trp Leu Ile Ala Val His Ser Ser Ile Phe Ser Ser Val Ala Ala Ser Cys Thr Leu Phe Val Val Leu Trp Leu Arg Val Pro Met Leu Arg Thr 225 235 240 Nal Phe Gly Phe Arg Trp Leu Gly Ala Ile Phe Leu Ser Asn Ser Arg 245 250

### (2) INFORMATION FOR SEQ ID NO:68:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 765 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: unknown
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: cDNA
- (vi) ORIGINAL SOURCE:
  - (A) ORGANISM: porcine reproductive and respiratory syndrome virus
  - (B) STRAIN: Iowa
  - (C) INDIVIDUAL ISOLATE: ISU-12 (VR 2385/VR 2386)
- (ix) FEATURE:
  - (A) NAME/KEY: CDS

## (B) LOCATION: 1..762

# (xi) SEQUENCE DESCRIPTION: SEQ ID NO:68:

	,,							-			-						
													TGC Cys				48
TTG Leu	TAC Tyr	TCT Ser	TTT Phe 20	TGT Cys	TGT Cys	GCT Ala	GTG Val	GTT Val 25	GCG Ala	GGT Gly	TCC Ser	AAT Asn	GCT Ala 30	ACG Thr	TAC Tyr		96
															ACG Thr		144
GTG Yal	AAT Asn 50	TAC Tyr	ACG Thr	GTG Val	TGC Cys	CCG Pro 55	CCT Pro	TGC Cys	CTC Leu	ACC Thr	CGG Arg 60	Gln	GCA Ala	GCC Ala	GCA Ala		192
											Arg		GGG Gly		GAT Asp 80	,	240
CGA Arg	TGT Cys	GGG Gly	GAG Glu	GAC Asp 85	GAT Asp	CAT His	GAT Asp	GAA Glu	CTA Leu 90	Gly	TTT Phe	GTG Val	GTG Val	CCG Pro 95	Ser		288
				Glu					Ser				TGG Trp 110	Leu	GCG Ala		336
			Phe					Gln							GGG Gly		384
		Asn					Tyr					His	CAA Glr		ATT : Ile		432
Cys		Val	His	Asp		Gln	Asr	1 Thr	Thr	Leu	Pro	Hi	CAT s His	a Asr	AAC Asn 160		480
					Gln					His			GAC l Asp		g Gly		528
				Leu					Pro				TCT r Set 190	r Trp	TTG Leu		576

								AGG Arg									624
								AGA Arg									672
								GTT Val									720
								CTC Leu									762
TAG			•			٠											765
⊕ (2)	INFO	ORMA'	rion	FOR	SEQ	ID I	NO:6	9:									
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i	(:	Li) 1			POLO:												
			MOLE	CULE	TYP	E: p:	rote		Q ID	NO:	69:		•				
	(2	ki) s	MOLE SEQUI	CULE ENCE	TYP	E: p: CRIP'	rote TION	in : SE(	-			Cys	Cys	Ser 15	Phe		
Met 1	(z Ala	ki) : Asn	MOLE SEQUI Ser	CULE ENCE Cys 5	TYP: DES	E: p: CRIP' Phe	rote FION Leu	in : SE(	Ile 10	Phe	Leu			. 15			
Met 1 Leu	() Ala Tyr	ki) s Asn Ser	MOLE SEQUI Ser Phe 20	CULE ENCE Cys 5 Cys	TYP: DESC Thr Cys	E: p: CRIP Phe Ala	rote FION Leu Val	in : SEG Tyr Val	Ile 10 Ala	Phe	Leu	Asn	Ala 30	Thr	Tyr		
Met 1 Leu	(z Ala Tyr Phe	Asn Ser Trp 35	SEQUI Ser Phe 20 Phe	CULE ENCE Cys 5 Cys	TYPE DESCRIPTION Thr Cys Leu	E: p: CRIP Phe Ala Val	rote FION Leu Val Arg 40	in : SEG Tyr Val 25 Gly	Ile 10 Ala Asn	Phe Gly Phe	Leu Ser Ser	Asn Phe 45	Ala 30 Glu	Thr Leu	Tyr		
Leu Cys	Ala Tyr Phe Asn 50	Asn Ser Trp 35	MOLEO SEQUI Ser Phe 20 Phe Thr	CULE ENCE Cys 5 Cys Pro Val	TYP: DESC Thr Cys Leu Cys	E: p: CRIP Phe Ala Val Pro 55 Arg	rote FION Leu Val Arg 40 Pro	in : SEG Tyr Val 25 Gly	Ile 10 Ala Asn Leu	Phe Gly Phe Thr	Leu Ser Ser Arg	Asn Phe 45 Gln	Ala 30 Glu Ala	Thr Leu Ala	Tyr Thr Ala		
Leu Cys Val Glu 65	Ala Tyr Phe Asn 50 Ala	Asn Ser Trp 35 Tyr	SEQUI Ser Phe 20 Phe Thr	CULE ENCE Cys Cys Pro Val	TYPE DESCRIPTION Cys Leu Cys Gly 70 Asp	E: p: CRIP Phe Ala Val Pro 55 Arg	rote FION Leu Val Arg 40 Pro	in : SEG Tyr Val 25 Gly Cys	Ile 10 Ala Asn Leu	Phe Gly Phe Thr Cys 75	Leu Ser Ser Arg 60	Asn Phe 45 Gln	Ala 30 Glu Ala Gly	Thr Leu Ala	Tyr Thr Ala Asp 80 Ser		

Ser Leu Ser Phe Ser Tyr Thr Ala Gln Phe His Pro Glu Ile Phe Gly 115 Ile Gly Asn Val Ser Arg Val Tyr Val Asp Ile Lys His Gln Phe Ile 135 Cys Ala Val His Asp Gly Gln Asn Thr Thr Leu Pro His His Asp Asn Ile Ser Ala Val Leu Gln Thr Tyr Tyr Gln His Gln Val Asp Gly Gly Asn Trp Phe His Leu Glu Trp Val Arg Pro Phe Phe Ser Ser Trp Leu Val Leu Asn Val Ser Trp Phe Leu Arg Arg Ser Pro Ala Ser His Val Ser Val Arg Val Phe Gln Thr Ser Arg Pro Thr Pro Pro Gln Arg Gln 215 ` Ala Leu Leu Ser Ser Lys Thr Ser Val Ala Leu Gly Ile Ala Thr Arg 225 235 HPro Leu Arg Arg Phe Ala Lys Ser Leu Ser Ala Ala Arg Arg 250 (2) INFORMATION FOR SEQ ID NO:70:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 537 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: unknown
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: cDNA
- (vi) ORIGINAL SOURCE:
  - (A) ORGANISM: porcine reproductive and respiratory syndrome virus
  - (B) STRAIN: Iowa
  - (C) INDIVIDUAL ISOLATE: ISU-12 (VR 2385/VR 2386)
- (ix) FEATURE:

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- (A) NAME/KEY: CDS
- (B) LOCATION: 1..534

### (xi) SEQUENCE DESCRIPTION: SEQ/ID NO:70: 48 ATG GGT GCG TCC CTT CTT TTC CTC TTG GTT GGT TTT AAA TGT CTC TTG Met Gly Ala Ser Leu Leu Phe Leu Leu Val Gly Phe Lys Cys Leu Leu GTT TCT CAG GCG TTC GCC TGC AAG CCA TGT TTC AGT TCG AGT CTT TCA 96 Val Ser Gln Ala Phe Ala Cys Lys Pro Cys Phe Ser Ser Leu Ser GAC ATC AAG ACC AAC ACC GCA GCG GCA GGC TTT GCT GTC CTC CAA 144 Asp Ile Lys Thr Asn Thr Thr Ala Ala Ala Gly Phe Ala Val Leu Gln GAC ATC AGT TGC CTT AGG CAT CGC AAC TCG GCC TCT GAG GCG ATT CGC 192 Asp Ile Ser Cys Leu Arg His Arg Asn Ser Ala Ser Glu Ala Ile Arg 50 AAA GTC CCT CAG TGC CGC ACG GCG ATA GGG ACA CCC GTG TAT ATC ACT 240 Lys Val Pro Gln Cys Arg Thr Ala Ile Gly Thr Pro Val Tyr Ile Thr \_\_\_65 GTC ACA GCC AAT GTT ACC GAT GAG AAT TAT TTG CAT TCC TCT GAT CTT 288 ∜al Thr Ala Asn Val Thr Asp Glu Asn Tyr Leu His Ser Ser Asp Leu 90 95 85 TC ATG CTT TCT TCT TGC CTT TTC TAT GCT TCT GAG ATG AGT GAA AAG 336 Leu Met Leu Ser Ser Cys Leu Phe Tyr Ala Ser Glu Met Ser Glu Lys 100 TGGA TTT AAG GTG GTA TTT GGC AAT GTG TCA GGC ATC GTG GCA GTG TGC 384 Gly Phe Lys Val Val Phe Gly Asn Val Ser Gly Ile Val Ala Val Cys 115 GTC AAC TTC ACC AGT TAC GTC CAA CAT GTC AAG GAA TTT ACC CAA CGT 432 Val Asn Phe Thr Ser Tyr Val Gln His Val Lys Glu Phe Thr Gln Arg 130 TCC TTG GTA GTT GAC CAT GTG CGG CTG CTC CAT TTC ATG ACG CCC GAG 480 Ser Leu Val Val Asp His Val Arg Leu Leu His Phe Met Thr Pro Glu 155 145 150 ACC ATG AGG TGG GCA ACT GTT TTA GCC TGT CTT TTT ACC ATT CTG TTG 528

Thr Met Arg Trp Ala Thr Val Leu Ala Cys Leu Phe Thr Ile Leu Leu

165

GCA ATT TGA Ala Ile 170

175

537

- (2) INFORMATION FOR SEQ ID NO:71:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 178 amino acids
    - (B) TYPE: amino acid
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: protein
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:71:

Met Gly Ala Ser Leu Leu Phe Leu Leu Val Gly Phe Lys Cys Leu Leu 1 5 10 15

Val Ser Gln Ala Phe Ala Cys Lys Pro Cys Phe Ser Ser Ser Leu Ser 20 25 30

Asp Ile Lys Thr Asn Thr Thr Ala Ala Ala Gly Phe Ala Val Leu Gln
45

Asp Ile Ser Cys Leu Arg His Arg Asn Ser Ala Ser Glu Ala Ile Arg
50 55 60

Lys Val Pro Gln Cys Arg Thr Ala Ile Gly Thr Pro Val Tyr Ile Thr
5 70 75 80

Val Thr Ala Asn Val Thr Asp Glu Asn Tyr Leu His Ser Ser Asp Leu
85 90 95

□Leu Met Leu Ser Ser Cys Leu Phe Tyr Ala Ser Glu Met Ser Glu Lys
□ 100 105 110

Gly Phe Lys Val Val Phe Gly Asn Val Ser Gly Ile Val Ala Val Cys 115 120 125

Val Asn Phe Thr Ser Tyr Val Gln His Val Lys Glu Phe Thr Gln Arg 130 135 140

Ser Leu Val Val Asp His Val Arg Leu Leu His Phe Met Thr Pro Glu 145 150 155 160

Thr Met Arg Trp Ala Thr Val Leu Ala Cys Leu Phe Thr Ile Leu Leu 165 170 175

Ala Ile

- (2) INFORMATION FOR SEQ ID NO:72:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 750 base pairs
    - (B) TYPE: nucleic acid

- (C) STRANDEDNESS: unknown
- (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: cDNA
- (vi) ORIGINAL SOURCE:
  - (A) ORGANISM: porcine reproductive and respiratory syndrome

virus

- (C) INDIVIDUAL ISOLATE: Lelystad
- (ix) FEATURE:
  - (A) NAME/KEY: CDS
  - (B) LOCATION: 1..747
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:72:

ATG Met	CAA Gln	TGG Trp	GGT Gly	CAC His 5	TGT Cys	GGA Gly	GTA Val	Lys AAA	TCA Ser 10	GCC Ala	AGC Ser	TGT Cys	TCG Ser	TGG Trp 15	ACG Thr	48
CCT Pro	TCA Ser	CTG Leu	AGT Ser 20	TCC Ser	TTG Leu	TTA Leu	GTG Val	TGG Trp 25	TTG Leu	ATA Ile	TTG Leu	CCA Pro	TTT Phe 30	TCC Ser	TTG Leu	96
CCA	TAC Tyr	TGT Cys 35	TTG Leu	GGT Gly	TCA Ser	CCG Pro	TCG Ser 40	CAG Gln	GAT Asp	GGT Gly	TAC Tyr	TGG Trp 45	Ser	TTC Phe	TTC Phe	144
TCA	GAG Glu 50	TGG Trp	TTT Phe	GCT Ala	CCG Pro	CGC Arg 55	TTC Phe	TCC Ser	GTT Val	CGC Arg	GCT Ala 60	Leu	CCA Pro	TTC Phe	ACT Thr	192
CTC Leu 65	CCG Pro	AAC Asn	TAT Tyr	CGA Arg	AGG Arg 70	TCC Ser	TAT	GAA Glu	GGC Gly	TTG Leu 75	Lev	CCC Pro	AAC Asn	TGC Cys	AGA Arg 80	240
										Pro			ATG Met		Trp	288
CAC His	ATG Met	CGA Arg	GTT Val 100	Ser	CAC His	TTG Leu	ATT Ile	GAT Asp 105	Glu	ATG Met	GTC Val	TCT Sei	CGT Arg	Arg	ATT J Ile	336
TAC Tyr	CAG Gln	ACC Thr 115	Met	GAA Glu	CAT His	TCA Ser	GGT Gly 120	Gln	GCG Ala	GCC Ala	TGG Trj	AAG Lys 12!	в Glr	GTG val	GTT L Val	384
GGT Gly	GAG Glu 130	Ala	ACT Thr	CTC Leu	ACG Thr	AAG Lys 135	Lev	TCA Ser	GGG Gly	CTC Let	GAT Asj 14	p Ile	GTT e Val	ACT L Thi	CAT r His	432

													TTT (			480
													GTG . Val			528
													CCC Pro 190			576
								Phe					ATC . Ile			624
CAC His							Val					Thr			ATA Ile	672
GTG Val 225											Val					720
TGG Trp																750
[≟ .ౖ(2)	INF	ORMA	TION	FOR	SEQ	ID	NO:7	'3:			•			٠		
		(i)	(A (B	ENCE ) LE ) TY ) TO	NGTH PE:	: 24 amin	9 am	ino id	: acid	ls					· :	
	(	ii)	MOLE	CULE	TYP	E: p	rote	ein								
	(	xi)	SEQU	ENCE	DES	CRIE	OIT	I: SE	Q II	NO:	73 :					
Met 1		Trp	Gly	His 5	Cys	Gly	⁄ Val	Lys	Ser 10		a Sei	c Cys	s Ser	Trp 15	Thr	
Pro	Ser	Leu	Ser 20		Leu	ı Lev	ı Val	Trp 25	-	ı Ile	e Lei	ı Pro	Phe 30		Leu	
Pro	Tyr	Cys 35		Gly	ser Ser	Pro	Ser 40	_	n Asp	Gly	у Ту:	r Try 49	_	Phe	Phe	

Ser Glu Trp Phe Ala Pro Arg Phe Ser Val Arg Ala Leu Pro Phe Thr 50 55 60

Leu Pro Asn Tyr Arg Arg Ser Tyr Glu Gly Leu Leu Pro Asn Cys Arg Pro Asp Val Pro Gln Phe Ala Val Lys His Pro Leu Gly Met Phe Trp His Met Arg Val Ser His Leu Ile Asp Glu Met Val Ser Arg Arg Ile 100 105 Tyr Gln Thr Met Glu His Ser Gly Gln Ala Ala Trp Lys Gln Val Val Gly Glu Ala Thr Leu Thr Lys Leu Ser Gly Leu Asp Ile Val Thr His 130 Phe Gln His Leu Ala Ala Val Glu Ala Asp Ser Cys Arg Phe Leu Ser Ser Arg Leu Val Met Leu Lys Asn Leu Ala Val Gly Asn Val Ser Leu Gln Tyr Asn Thr Thr Leu Asp Arg Val Glu Leu Ile Phe Pro Thr Pro 180 185 Gly Thr Arg Pro Lys Leu Thr Asp Phe Arg Gln Trp Leu Ile Ser Val His Ala Ser Ile Phe Ser Ser Val Ala Ser Ser Val Thr Leu Phe Ile 210 Val Leu Trp Leu Arg Ile Pro Ala Leu Arg Tyr Val Phe Gly Phe His \_225 230 235 Trp Pro Thr Ala Thr His His Ser Ser

(2) INFORMATION FOR SEQ ID NO:74:

245

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 798 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: unknown
- (D) TOPOLOGY: linear

### (ii) MOLECULE TYPE: cDNA

#### (vi) ORIGINAL SOURCE:

- (A) ORGANISM: porcine reproductive and respiratory syndrome virus
- (C) INDIVIDUAL ISOLATE: Lelystad



(ix) FEATURE:
(A) NAME/KEY: CDS
(B) LOCATION: 1..795

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:74:

	(xi)	SEÇ	QUENC	CE DI	ESCR	[PTIC	ON:	SEQ :	ID No	0:74	:						
			CAG Gln														48
			GTT Val 20														96
			TTT Phe										Glu				144
ATC Ile			ACC Thr									Gln					192
CAA Gln 65			GAG Glu								Lys						240
AGG Arg					Asp					Leu					Ser		288
GGG -Gly									Tyr					Ala		مدائيت والعدي	336
								Phe					ı Phe		ATA / Ile		384
			TCG Ser				٧al					Glr			TGT TGT	·	432
	Glu		GAT Asp			Asn					Thr				ATC lle 160		480
			TAT Tyr		Ala					Glr					/ Asn		528

			TTG Leu 180											Leu		57	6
			TCA Ser													62	4
			TAT Tyr													67	12
			TTC Phe								Leu				CAG Gln 240	72	20
CAG -Gln			AGA Arg							Arg					Lys	7€	8
Pro										•						79	8
(2)	INF	ORMA	rion	FOR	SEQ	·ID :	NO : 7	5:					,		, .		
(2) INFORMATION FOR SEQ ID NO:75:  (i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 265 amino acids  (B) TYPE: amino acid  (D) TOPOLOGY: linear  (ii) MOLECULE TYPE: protein																	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:75:																
Met 1										Phe		Cys	Gly	Phe 15	Ile		
Cys	Tyr	Leu	Val 20	His	Ser	Ala	Leu	Ala 25		Asn	Ser	Ser	Ser 30		Leu		
Сув	Phe	Trp 35	Phe	Pro	Leu	Ala	His 40	-	Asn	Thr	Ser	Phe 45		ı Leu	Thr		
Ile	Asn 50		Thr	Ile	Cys	Met 55		Cys	Ser	Thr	Ser 60		a Ala	a Ala	Arg		

Gln Arg Leu Glu Pro Gly Arg Asn Met Trp Cys Lys Ile Gly His Asp 65 70 75 80

Arg Cys Glu Glu Arg Asp His Asp Glu Leu Leu Met Ser Ile Pro Ser Gly Tyr Asp Asn Leu Lys Leu Glu Gly Tyr Tyr Ala Trp Leu Ala Phe Leu Ser Phe Ser Tyr Ala Ala Gln Phe His Pro Glu Leu Phe Gly Ile Gly Asn Val Ser Arg Val Phe Val Asp Lys Arg His Gln Phe Ile Cys Ala Glu His Asp Gly His Asn Ser Thr Val Ser Thr Gly His Asn Ile 145 150 155 160 Ser Ala Leu Tyr Ala Ala Tyr Tyr His His Gln Ile Asp Gly Gly Asn 170 Trp Phe His Leu Glu Trp Leu Arg Pro Leu Phe Ser Ser Trp Leu Val 180 185 Leu Asn Ile Ser Trp Phe Leu Arg Arg Ser Pro Val Ser Pro Val Ser M 195 200 Arg Arg Ile Tyr Gln Ile Leu Arg Pro Thr Arg Pro Arg Leu Pro Val Ser Trp Ser Phe Arg Thr Ser Ile Val Ser Asp Leu Thr Gly Ser Gln **2225** 230 235 Gln Arg Lys Arg Lys Phe Pro Ser Glu Ser Arg Pro Asn Val Val Lys 250 ]= Pro Ser Val Leu Pro Ser Thr Ser Arg 260 265

#### (2) INFORMATION FOR SEQ ID NO:76:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 552 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: unknown
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: cDNA
- (vi) ORIGINAL SOURCE:
  - (C) INDIVIDUAL ISOLATE: Lelystad
- (ix) FEATURE:
  - (A) NAME/KEY: CDS
  - (B) LOCATION: 1..549



# (xi) SEQUENCE DESCRIPTION: SEQ ID NO:76:

	(XI)	وعدد	SORIAC		ocic.	LFII(	J14	JEQ .	ID IV	J. 76	•						
													CAT His			4	48
													CAT His 30			, <b>!</b>	96
															CAG Gln	14	44
													CAA Gln			19	92
ATT LILE 65																2	40
CAG Gln																2	88
													GCC Ala 110	Ser	GAA Glu	3	36
ATG Met								Ile					Ser				84
												His	GTG Val		CAA Gln	4	32
	Thr			His		Leu			Asp		Ile		TTG Leu		CAT His 160	<b>.</b> . <b>4</b>	80
										Thr			GCT Ala		Leu	5	28
						ATA Ile										5	52



#### (2) INFORMATION FOR SEQ ID NO:77:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 183 amino acids
  - (B) TYPE: amino acid
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:77:

Met Ala Ala Thr Leu Phe Phe Leu Ala Gly Ala Gln His Ile Met
1 5 10 15

Val Ser Glu Ala Phe Ala Cys Lys Pro Cys Phe Ser Thr His Leu Ser 20 25 30

Asp Ile Glu Thr Asn Thr Thr Ala Ala Ala Gly Phe Met Val Leu Gln
45

Asp Ile Asn Cys Phe Arg Pro His Gly Val Ser Ala Ala Gln Glu Lys
50 55 60

Tle Ser Phe Gly Lys Ser Ser Gln Cys Arg Glu Ala Val Gly Thr Pro 65 70 75 80

Gln Tyr Ile Thr Ile Thr Ala Asn Val Thr Asp Glu Ser Tyr Leu Tyr
90
95

Asn Ala Asp Leu Leu Met Leu Ser Ala Cys Leu Phe Tyr Ala Ser Glu

Met Ser Glu Lys Gly Phe Lys Val Ile Phe Gly Asn Val Ser Gly Val 115 120 125

Val Ser Ala Cys Val Asn Phe Thr Asp Tyr Val Ala His Val Thr Gln 130 135 140

His Thr Gln Gln His His Leu Val Ile Asp His Ile Arg Leu Leu His 145 155 160

Phe Leu Thr Pro Ser Ala Met Arg Trp Ala Thr Thr Ile Ala Cys Leu 165 170 175

Phe Ala Ile Leu Leu Ala Ile 180